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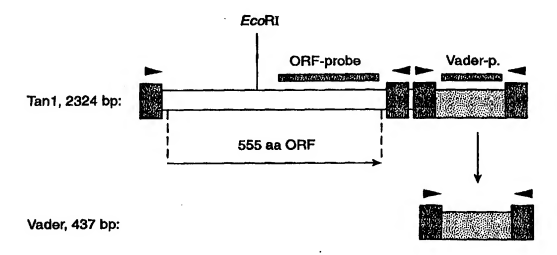
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(54) Title: IDENTIFICATION OF AND CLONING A MOBILE TRANSPOSON FROM ASPERGILLUS



(57) Abstract

There are provided transposable elements isolated from Aspergillus. Also provided are fragments comprising the inverted repeat(s) of the transposable elements, such fragments being useful as probes to isolate transposable elements from other filamentous fungi.

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IDENTIFICATION OF AND CLONING A MOBILE TRANSPOSON FROM ASPERGILLUS

Field of the Invention

The present invention is directed at the identification, cloning and sequencing of mobile transposons or transposable elements from *Aspergillus niger* var. *awamori*. The transposable elements, referred to as Vader and Tan1, are approximately 437 base pair (bp) and 2.3 kb elements, respectively. The Vader and Tan1 elements are bounded by inverted repeat sequences of 44 and 45 base pairs, respectively. The transposable elements target a "TA" sequence in target DNA during insertion. In addition, the present invention is directed at the identification, cloning and sequencing of one or more transposable element(s) from other filamentous fungi using as a probe DNA comprising the Vader element 44 bp or the Tan1 element 45 bp inverted repeat isolated from *Aspergillus niger* var. *awamori*. Also provided are methods for utilizing either the Vader or Tan1 elements to inactivate genes (for example, by inserting the transposon into the gene to be inactivated), to overexpress a gene (by, for example, inserting a known promoter or other regulatory gene within the inverted repeats of Vader or Tan 1 and allowing the DNA of the IR-promoter-IR to jump in front of (and overexpress) a gene of interest) or to act as an activation marker to, for example, identify new promoters.

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Background of the Invention

It is well know that transposons are a class of DNA sequences that can move from an episome to a chromosomal site or from one chromosomal site to another. Transposons are known in both prokaryotes, such as bacteria, as well as in eukaryotes, although there have been few transposons isolated from filamentous fungi.

Several groups have looked for transposons in filamentous fungi. The element pogo, which exists in multiple copies and at different sites in different strains of Neurospora crassa, was described by Schectman (1) and is believed to be a transposon. To date the most characterized transposon in filamentous fungi is Tad. Tad was isolated as a spontaneous mutant in the am (glutamate dehydrogenase) gene in an Adiopodoume strain of N. crassa isolated from the Ivory Coast. To detect mutations caused by insertion of a transposable element, Kinsey and Helber (2) isolated genomic DNA from 33 am mutant

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transposable element, Kinsey and Helber (2) isolated genomic DNA from 33 am mutant strains which were then screened by Southern analysis for restriction fragment size alterations. In two of the mutant strains, the mutation was shown to be caused by the insertion of a 7 kb element (Tad) into the am gene. Subsequently Kinsey (3) demonstrated that Tad was able to transpose between nuclei of heterokaryons, confirming that Tad was a retrotransposon and that there was a cytoplasmic phase involved in the retrotransposition events. More recently, Cambareri et al. (4) demonstrated that Tad was a LINE-like DNA element with two major open reading frames (ORFs) on the plus strand. Typical of LINE-like elements, Tad had no terminal repeats. Attempts to isolate mobile transposons in laboratory strains of *N. crassa* were unsuccessful.

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A second retrotransposon was cloned by McHale et al. (5), who reported the isolation of CfT-1, an LTR-retrotransposon from *Cladosporium fulvum*. This transposon was 6968 bp in length and bounded by identical long terminal repeats of 427 bp, a 5 bp target site duplication. Virus-like particles were detected which co-sediment with reverse transcriptase activity in homogenates of this fungus.

Daboussi et al. (6) were the first to successfully use the niaD (nitrate reductase) gene as a transposon trap. The *niaD* mutants can be isolated by a direct selection for chlorate resistance (7). The strategy employed was to isolate niaD mutants amongst six isolates belonging to different races of the fungus Fusarium oxysporum. More than 100 niaD mutants were isolated from each isolate and examined for instability. One strain, F24, vielded up to 10% unstable niaD mutants. Assuming that the genetic instability of the niaD mutants was caused by transposable elements, it seemed plausible that this isolate contained mobile transposons. A stable niaD mutant in the F24 was transformed with the cloned niaD gene from A. nidulans because the F. oxysporum niaD gene had not been cloned. Unstable niaD mutants were isolated in transformants containing the A. nidulans niaD gene. Two unstable niaD mutants were shown by Southern blot analysis to contain a insertion of 1.9 kb in size. Analysis of this element, Fot1, revealed it was 1928 bp long, had a 44 bp inverted terminal repeats, contained a large open reading frame, and was flanked by a 2 bp (TA) target site duplication. Very recently, Daboussi et al. (8) have reported the cloning of a new transposable element from an unstable niaD mutant. This element, FML (Fusarium mariner-like), is 1280 bp long and has inverted repeats of 27 bp. The FML element inserts into a TA site and excises imprecisely.

Using the characterization of unstable *niaD* mutants strategy, Lebrun et al. (9) were able to isolate a transposon from *Magnaporthe grisea*. However, in this case the A.

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nidulans niaD gene which was transformed into *M. grisea* by transformation was used as a transposon trap. The element inserted into the *niaD* gene was shown to belong to a family of *M. grisea* LTR-retrotransposons, Fos1 (Schull and Hamer, unpublished) and Mag1 (Farman and Leong, unpublished). The cloned retro-element was 5.6 kb and the target site (ATATT) was shown to be duplicated. All revertants from this mutant examined had one copy of the LTR left at the point of insertion. A second transposon, Pot2, from *M. grisea* was recently cloned by Kachroo et al. (10). The strategy used to clone Pot2 was to analyze the fingerprint patterns of repetitive DNA's which were cloned from the *M. grisea* genome. A repetitive family present in both rice and non-rice pathogens of *M. grisea* in high copy number was cloned. The element, 1857 bp in size, has a 43 bp perfect terminal inverted repeats (TIR) and 16 bp direct repeats within the TIRs. An open reading frame was shown to display extensive identity to that of Fot1 of *F. oxysporum*. As with Fot1, the Pot2 element duplicates the dinucleotide TA at the target insertion site. Pot2 was shown to be present at a copy number of approximately 100 per haploid genome.

Several groups have reported looking without success for transposons in laboratory strains of *A. nidulans* (Kinghorn personnel communication, 5). One explanation for the lack of transposons in laboratory strains is that the desirable features of strain stability required for genetic analysis may preclude strains with mobile transposon. By using the *niaD* gene as a transposon trap we have identified and isolated a transposable element from the industrially important fungus *A. niger* var. *awamoni*. This element, Vader, is present in approximately 15 copies in *A. niger* and *A. niger* var. *awamoni*. Southern analysis of *A. nidulans* with this element indicates that this transposable element was absent from one laboratory strain and only present as a single copy in a second laboratory strain. These results support the notion that laboratory strains of *A. nidulans* contain very few transposons.

Brief Description of the Invention

In accordance with the present invention, novel eukaryotic transposable elements from Aspergillus niger var. awamori are provided. The larger transposable element, referred to herein as Tan1, is 2.3 kb in size. The smaller transposable element, referred to herein as Vader, is a 437 bp element (SEQ ID NO:3). Vader is found within the larger element Tan1. The Vader transposable element is a 437 bp element which comprises a 44 bp inverted repeat sequence at either end of the transposable element. Tan1 is approximately a 2325 bp element which comprises 45 bp inverted repeats at either terminus and internal IRs. Tan1 comprises a 555 aa open reading frame (ORF) which

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codes for a transposase which allows the elements (Tan1 or Vader) to "hop" or insert themselves in the genome of a host. The target for insertion of these novel transposable elements is a "TA" sequence in the target DNA for insertion. The "TA" sequence is repeated at either end of the transposon upon insertion of the transposable element into the target DNA. Therefore, the present invention provides the larger Tan1 transposable element as well as the smaller element (Vader) internal thereto, as well as the DNA encoding each.

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Another embodiment of the present invention comprises a fragment of the Vader or Tan1 transposable elements which comprise the 44 or 45 bp (respectively) inverted repeat sequences found at either terminus of the transposable element from *A. niger* var. awamori, as well as the use of said fragments as probes to hybridize under low stringency conditions to DNA of other filamentous fungi for the isolating and/or cloning of transposable elements from such other filamentous fungi. While the exact 44 bp IR of Vader or the 45 bp IR of Tan1 can be utilized, it is well understood by those skilled in the art that variation of such DNA would also work as a suitable probe. For example, at a minimum, the imperfect direct repeats within the IRs of Tan1 would be suitable to use as probes for isolating transposable elements from other filamentous fungi. Initially the inverted repeat of Vader was used to clone Tan1 using PCR techniques. This work was followed by obtaining a genomic copy of Tan1 from a partial library.

Another embodiment of the present invention is the transposase activity coded for by the ORF of Tan1. This transposase is 555 aa (SEQ ID NOS:7 or 14, PCR and genomic, respectively).

In a process embodiment of the present invention there are provided methods for gene tagging comprising using the transposable elements of the present invention (Vader or Tan1 or any transposable element isolated using the IRs of either) to inactivate genes via insertion of the element into a given gene, thus disrupting or inactivating gene expression. Alternatively, the transposable element can be used in activation tagging (to activate or turn on genes) rather than for gene disruption. For example, by inserting DNA coding a promoter into the transposable element and then allowing such transposable element to become inserted 5' to a desired gene, the promoter may be activated to drive the expression of the desired gene product or to turn on cryptic pathways. Additionally, gene tagging can be utilized to activate marker genes by inserting a marker gene within the IRs of a transposon of the present invention. This marker gene can then "hop" into targeted DNA and, if expression of the marker is selected for, it will be possible to identify

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the promoter driving such expression. This may lead to identification of isolation of new strong promoters.

Brief Description of the Drawings

Fig. 1 shows the Southern blot analysis of unstable *niaD* mutants. PCR-amplified genomic *niaD* gene from four *niaD* mutants and UVK143f were digested with Bglll (sites are 3' of all inserts). Blot probed with 500 bp fragment of Sall digested PCR product of niaD1 and niaD2. Wild-type band hybridizes at 2.5 kb while gene with insertion hybridizes at 2.9 kb. Lanes: 1=MW marker III (Boehringer Mannheim); 2=UVK143f; 3=*niaD*410; 4=*niaD*436; 5=*niaD* 587; 6=*niaD*392.

Fig. 2 depicts the mapping of Vader insertions within the *niaD* gene. The positions of Vader insertions 1-4 (*nia*D410, *nia*D436, *nia*D587 and *nia*D392, respectively) are shown relative to the six introns of the structural gene coding region. Because the exact site of insertion for Vader-1 and Vader-4 is still unknown, they have been presented using the approximate area of insertion. Relevant restriction sites are shown using the following letters: E=EcoRI, S=SalI, Sp=SphI, K=KpnI, and B=BgIII.

Fig. 3 shows Southern blot analysis to determine Vader genomic copy number. Four *A. niger* var. *awamori niaD* mutants and UVK143f were digested with EcoRV to completion. EcoRV cuts the Vader sequence once. Hybridization indicates that Vader is present in the genome in more than 14 copies. The hybridizing bands of *niaD* 392, which are different from the other mutants and UVK143f, suggest that the Vader sequence is mobile. Lanes: 1=MW marker III, 2=UVK143f, 3=*niaD*410, 4=*niaD*436, 5=*niaD*587, 6=*niaD*392.

- Fig. 4. Southern blot to determine presence of Vader sequence in other fungi.

 Other filamentous fungi, an industrial production strain and *niaD* mutant 392 were digested with EcoRV to completion. Low stringency hybridization (32) indicates that sequences homologous to Vader are present in *A. nidulans* (FGSC A237), *A. cinnamomeus*, *A. phoenicis*, *A. foetidus*, an industrial *A. niger* strain. Lanes: 1=MW marker, 2=*A. foetidus*, 3=an industrial glucoamylase production strain of *A. niger* (ETC #2663), 4=*A. niger* var. awamori niaD mutant 392, 5=*A. phoenicis* (ATCC #11362), 6=*A. nidulans* (FGSC A691), 7=*A. wentii* (ATCC #10593), 8=*A. versicolor*, 9= *A. cinnamomeus* (ATCC #1027), 10=*A. nidulans* (FGSC A237)
- Fig. 5. Southern blot to determine Tan1 (transposon from A. niger) genomic copy number. Four niaD mutants A. niger var. awamori mutants and UVK143f were digested

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with *Eco*RI to completion. *Eco*RI cuts the Tan1 sequence once. A probe corresponding to the ORF region (see Fig. 9) was used in the hybridization. Hybridization indicates that Tan1 is present as a single copy in the genome. Lanes: 1=MW marker III, 2=UVK143f, 3=niaD410, 4=niaD 436, 5=niaD 587, 6=niaD 392.

Figs. 6A-6C. Southern blots to determine if the inverted repeats of transposable elements Fot1 and Pot2 will hybridize to elements in *A. niger* var. *awamori*. Four *niaD* mutants *A. niger* var. *awamori* mutants were digested with *Eco*RI to completion. *Eco*RI cuts the Tan1 sequence once. Inverted repeat oligonucleotide probes of Vader (SEQ ID NO:5), Fot1 and Pot2 were labeled with digoxigenin (Boehringer Mannheim). Lanes: 1=MW marker III, 2=niaD436, 3=niaD587. Blot A (lanes 1-3) and B and C were probed with the labeled inverted repeat probes of Vader, Fot1 and Pot2, respectively.

Fig. 7 shows the sequence of the Vader insertion (SEQ ID NO:3) as generated by PCR. Vader was found to be 437 bp in length. The 44 bp inverted repeat of the Vader insert corresponding to SEQ ID NO:4 (the 5' IR) and SEQ ID NO:5 (the 3' IR), respectively, from the 5' end to the 3' end of Vader are underlined, the single mismatch which occurs in the inverted repeats is identified in bold, and the TA 2 bp duplication is shown in bold print. *niaD* sequences flanking the element are shown in lower case letters.

Figs. 8A and 8D show the entire DNA sequence of the Tan1 element (SEQ ID NO:6) as generated by PCR, as well as the putative amino acid sequence of the transposase coded for by Tan1 (SEQ ID NO:7). Tan1 as generated by PCR is 2320 bp in length (excluding the unknown nucleotides shown as "N" in the figure) and has a large open reading frame of 1668 bp which encodes for 555 amino acids (SEQ ID NO:7). Tan1 comprises the sequences of four inverted repeats (underlined) similar to those found in Vader.

Fig. 9 shows a schematic presentation of Vader and Tan1 elements. Dark boxes represent the 45 bp (Tan1) and 44 bp (Vader) inverted repeats. The unique *EcoRI* site in the Tan1 element was used for digestion of genomic DNA in Southern analysis (Figs. 5 and 10). Bold, horizontal lines above the Tan1 element indicate the probes corresponding to the end of the ORF and Vader used in Southern analysis shown in Fig. 10 and Fig. 5.

Fig. 10 shows Southern analysis of *A. niger* var. *awamon niaD* mutants (*niaD*410, *niaD*436, *niaD*587, *niaD*392) and the wild-type UVK143f: lane 1, molecular weight marker III (Boehringer Mannheim); lane 2, UVK143f; lane 3, *niaD*410; lane 4, *niaD*436; lane 5, *niaD*587; lane 6, *niaD*392. This blot was probed for the Vader element (see Fig. 9). When this blot (Fig. 10) was superimposed with the blot shown in Fig. 5, one of the illuminated bands from the Vader-probe hybridization overlaid the single band in the ORF-probe

hybridization indicating that the Tan1 element is composed of contiguous ORF and Vader elements.

Figs. 11A and 11D show the nucleotide sequence (genomic copy) of Tan1 (SEQ ID NO:13). The amino acid sequence encoding the putative transposase (555 aa) (SEQ ID NO:14) is shown below the DNA sequence in the one-letter amino acid code. The inverted repeats are underlined (SEQ ID NOS:1, 2, 15 and 16, respectively, 5' to 3') and the imperfect direct repeats within the inverted repeats are shown with arrows above or below the sequence. The gaps within the arrows indicate the imperfect nucleotides within the direct repeats. Undetermined sequence is denoted in the figure by question marks and in the sequence listing as "N." The figure shows the DNA sequence as 2324 base pairs, excluding the unknown nucleotides indicated by "?" in the figure.

Detailed Description of the Invention

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While the specification concludes with claims particularly pointing out and distinctly claiming the subject matter regarded as forming the present invention, it is believed that the invention will be better understood from the following detailed description of preferred embodiments.

Standard biochemical nomenclature is used herein in which the nucleotide bases are designated as adenine (A); thymine (T); guanine (G); and cytosine (C). N connotes any of these nucleotides. As is conventional for convenience in the structural representation of a DNA nucleotide sequence, only one strand is usually shown in which A on one strand connotes T on its complement and G connotes C.

Applicants have isolated two transposable elements from *A niger* var. *awamori*. The cloned element Vader was identified by screening unstable nitrate reductase (*niaD*) mutants for insertion. This element is present in approximately fifteen copies in the genome of *A. niger* strains examined. In contrast, the Vader element is present in one copy in only one of the two *A. nidulans* strains studied. These results explain why several groups have been unsuccessful in isolating active transposons in laboratory *A. nidulans* strains. A plausible assumption is that "domesticated" strains of *A. nidulans* have lost their transposons due to repeated manipulation of such strains and the possible discarding of aberrant *A. nidulans* strains displaying genetic instability.

The Vader element shows similarities to transposable elements cloned from the plant pathogens Pot1 from *M. grisea* (12) and Fot1 from *F. oxysporum* (8). The target site for duplication in all three fungi is a 2 bp TA sequence. In the case of Fot1, this transposon does not excise precisely. In two *niaD* revertants examined, the excision products retained a 4 bp insertion relative to the wild-type gene (TAATTA versus TA). The insertion studied

was integrated into an intron, therefore, imprecise excision of Fot1 did not effect the functionality of the *niaD* gene product. There is no published evidence that Pot2 is a functional element.

A homology search made at the nucleotide level gave a strong 60.7% homology between Tan1 and a 1230 bp overlap to the A. oryzae agdA gene coding for an α glucosidase (33). This homology search revealed that the last 1.2 kb of a total of 5.2 kb of the α -glucosidase sequence submitted to GenBank is, in fact, part of a novel transposon, hereinafter called Tao1 (transposon Aspergillus oryzae), which also belongs to the Fot1 family. Only the 5' half of the Tao1 element is included in the GenBank sequence, thus, for the lack of comparison, the exact size of the inverted repeat cannot be determined. However, it can be concluded that there are 13 bp perfect direct repeats within the inverted repeat. The inverted repeat is flanked by a TA-dinucleotide, suggesting a commonly occurring TA-insertion site. Direct analyses gave only short ORFs, but when the oftenoccurring stop codons were ignored, a long ORF was obtained which shared over 50% identity to the Tan1 transposase. Multiple stop codons indicate that the A. oryzae Tao1 is a defective element. This transposable element from A. oryzae, thus, is within the scope of the present invention as, based on the high degree of sequence homology between the Tan1 and Tao1, it is believed that Tao1 would hybridize to a probe comprising Tan1 or Vader IRs or variations thereof. The sequence of the IR of Tao1 is provided as SEQ ID NO:17. This IR (Tao1) or the IRs from Tan1 or Vader may be used to isolate other transposable elements from filamentous fungi.

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In an attempt to determine if there were transposons similar to those reported for *F. oxysporum* and *M. grisea*, synthetic oligomers were made corresponding to the inverted repeats of both Fot1 (7) and Pot2 (10). When Southern analysis of *A. niger* var. *awamoni* was conducted using the Vader 44 bp inverted repeat (SEQ ID NO:5) as a control, no conclusive hybridizations could be detected with either the Fot1 or Pot2 oligomeric probe. These results indicate that elements with high identity to *F. oxysporum* Fot1 and *M. grisea* Pot2 are not found from *A. niger* var. *awamoni* genome.

With regard to the structure of the Vader element, elements which transpose directly through DNA copies are typified by having inverted terminal repeats. Elements which transpose through reinsertion of the product of reverse transcription of an RNA copy of the element (retroelements) can be without long terminal repeats such as the *Drosphilia I* element (for a review see (16)). Alternatively, retrotransposons can have long terminal repeats such as the *Drosphilia copia* element. The Vader inverted repeats shown in Fig. 7,

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SEQ ID NOS:4 and 5, respectively, have a single mismatch. Elements which transpose through DNA copies typically have open reading frame(s) which encode a transposase activity. The Fot1 element is 1.9 kb in length and the Pot1 element 1.8 kb in length. Both the Fot1 and Pot1 elements have ORF encoding for a putative transposase-like protein. The Vader element, although mobile, does not have an ORF and hence it was deduced that the mobility of Vader was dependent upon a transposase activity present elsewhere in the genome. A synthetic 44 bp oligomer of the inverted repeat of Vader (SEQ ID NO:5) was used to clone, via PCR, a 2.3 kb element. This element, called Tan1 (SEQ ID NO:6), comprises four inverted repeats (SEQ ID NOS:1, 2, 15 and 16 from 5' to 3', respectively) similar to those in Vader and has a unique organization IR-ORF-IR-IR-Vader-IR. Tan1 is 2324 bp in length and has a large open reading frame (1668 bp) which encodes a putative transposase comprising 555 amino acids (shown in SEQ ID NOS:7 and 14), which is homologous to Fot1 and Pot2 transposases. Immediately 3' to the second IR (SEQ ID NO:2), which bounds the transposase, is a copy of the Vader element. We hypothesize that at some stage the independent Vader element, although inactive by itself, has arisen from Tan1, resulting in current strains with only one copy of Tan1 providing transposase activity and numerous mobile copies of Vader dispersed in the genome.

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Thus, applicants have been the first to identify a transposable element(s) with certain Aspergillus species. These transposable elements are believed to be quite useful in the development of gene tagging systems for Aspergillus or other microorganisms. Basic requirements for developing a gene tagging system are that the tagging element can be distinguished from the endogenous elements, it displays little sequence specificity for transposition and that excision is followed by integration at a new site. More refined tagging systems include ability to monitor excision and reinsertion by, e.g., activation of antibiotic resistance genes and ability to stabilize the mutations by, e.g., a two transposons system (23, 24 and 25).

For development of a tagging system for *Aspergillus*, it is proposed that the system is tested first in *A. nidulans*, which we have already shown does not have endogenous Tan1 or Vader sequences. However, at this stage the Vader element is altered from the original in such a way that the same construction can be later used in *A. niger* var. *awamori* and be distinguished from the endogenous Vader elements.

In a model tagging system using Vader as the "mutator," a first vector can be constructed for expression of the Vader element, similar to the non-autonomous maize Dc. The internal sequence of the Vader element is altered to contain translation initiation and stop codons in three different frames. This sequence can later be used as a recognition

site for a probe in PCR analysis of the mutants. This altered Vader element, Vader-S, is inserted within an expression cassette conferring antibiotic resistance such as hygromycin resistance. Since excision of Vader may not always be precise, Vader-S is inserted in the promoter area (e.g., oliC) between the transcription and translation initiation sites. This disrupted hygromycin phosphotransferase cassette is flanked by marker genes - or alternatively the marker gene upstream of the hygromycin promoter can be placed within Vader. These marker genes can be used for monitoring whether the hygromycin gene, and Vader within it, have integrated in full length. A vector, for example, Vector I, containing these elements will be transferred to A. nidulans and transformants expressing the two marker genes, but sensitive to hygromycin, are selected. Screening of mutants at later stages is easier, if the transformant selected for mutagenesis has only one to two copies of Vector I sequences integrated in its genome.

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A transformant with only a few (preferentially one) intact Vader-S/hygromycin phosphotransferase cassettes integrated in its genome is retransformed with Vector II, which is an autonomously replicating vector carrying the transposase encoding gene. The autonomously replicating vector, pHELP, used as a basis for DNA construction work, can be segregated away by methods known to those skilled in the art. This enables stabilization of the Vader-S element after the mutagenesis step. Vader-S is activated by a transposase (from Tan1) in pHELP, which can be monitored by activation of the hygromycin resistance gene. Tan1 is not cloned into the vector in full length to disrupt its mobility. Again, Vector II contains a marker gene used for screening of transformants and also for monitoring its segregation after the sporulation phase.

Marker genes can either complement host mutations or be dominant markers such as benomyl^R, acetamidase or β -glucuronidase (GUS).

In a model system for gene tagging the target gene for mutagenesis should be one with a simple plate screen, e.g., disruption of the *niaD* gene (by insertion of Vader), which can be screened by selection of chlorate resistant mutants and the gene disruption can be further mapped by a plate test using different nitrogen sources (no growth on nitrate, growth on nitrite, xanthine and uric acid). Another target gene for mutagenesis could be an acid protease gene. It has been shown previously for *A. niger* that disruption of this one protease is sufficient to abolish halo formation almost completely on skim milk plates.

The advantage of using transposon tagging is that the mutants produced can be identified by subsequent isolation of the mutated gene. There are several methods available for PCR amplification of genomic sequences when only one end of the sequence is known - which, in this case, is the transposable element. PCR methods developed for

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genomic walking are, e.g., "Inverse PCR" (27 and 28), "Vectorette PCR" (29) and "Panhandle PCR" (30).

Setting up the transposon tagging system can be followed by studies of excision frequency, environmental influences on transposition frequency (24, 31), activation of the transposase by a heterologous promoter and effect of altered inverted repeats on transposition.

Transposon tagging does need to be applied for inactivation of genes.

Alternatively, tagging can be used to insert promoter sequences in Vader and therein activate genes. A third option is to insert a promoterless marker gene in Vader, in which case the transposon can be used in search for novel, strong fungal promoters.

Experimental

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Materials and Methods

Strains. Vader and Tan1 elements were isolated from *Aspergillus niger* var. awamori UVK143f, derived from Northern Regional Research Laboratories (NRRL) #3112. *E. coli* JM101 [F' *traD36 lac1*^q Δ (*lacZ*)*M15 proA*[†]*B*[†] /supE thi Δ (*lac-proAB*)] and *Epicurian coli* SURE 2 (Stratagene Cloning Systems, La Jolla, CA) were used for propagation of Vader and Tan1 subclones, respectively.

Spontaneous chlorate resistant mutants were derived from Aspergillus niger var. awamori UVK143f (NRRL #3112). The following Aspergillus strains were obtained from the ATCC: A. cinnamomeus (ATCC #1027), A. wentii (ATCC #10593), and A. phoenicis (ATCC #11362). A. nidulans (FGSC #A237), a nitrate reductase structural gene mutant (niaD15), and A. nidulans (FGSC #A691), a tryptophan requiring mutant (trpC801), were obtained from Fungal Genetics Stock Center (FGSC), Dept. of Microbiology, University of Kansas Medical Center. A. versicolor, A. foetidus, and a proprietary A. niger glucoamylase strain are from the Genencor International Inc. culture collection.

Mutant Selection. Spore suspensions (1 x 10⁸) of UVK143f were plated on CM agar (11) containing 600 mM KClO₃ and 10 mM glutamic acid. Chlorate (KClO₃), a toxic analog of nitrate, allows selection of mutants in the nitrate assimilation pathway by chlorate resistance. Plates were incubated at 37°C until individual colonies of spontaneous mutants could be identified. Single mutants resistant to KClO₃ were allowed to sporulate on CM plates and spores from these plates were then streaked onto minimal media (11) with various sole nitrogen sources (10 mM): NaNO₃ (nitrate), NaNO₂ (nitrite), hypoxanthine, uric acid or NH₄Cl (ammonium chloride). Each of these compounds are intermediate products

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of the nitrate assimilation pathway. *niaD* mutants were identified as those resistant to KCIO₃ and able to grow in the presence of all pathway intermediates, except for NaNO₃.

Isolation of Vader via PCR Amplification. Genomic DNA of *A. niger* var. *awamori niaD* mutants and UVK143f was used as template (see Southern Analysis). Primers (50 pmol) used for amplification of the *niaD* gene were NiaD1 (position 142-165 relative to the initiation site of *niaD*): 5'-CCAACCGAGTCCTCAGTATAGAC-3' (SEQ ID NO:8) and NiaD2 (position 2738-2715): 5'-CAACGCTTCATAGGCGTCCAGATC-3' (SEQ ID NO:9). Deep Vent (exo⁻) DNA polymerase (New England Biolabs) was used with the buffer and dNTPs provided by the manufacturer. For optimal amplification of the *niaD* gene the reaction mixture contained 4 mM MgSO₄. Denaturation of template DNA, 2 min. at 94°C, was followed by 30 cycles of denaturation (30 sec. at 94°C), annealing of primers (45 sec. at 55°C) and extension (4 min. at 72°C). PCR fragments were purified from gel using the Qiaex DNA gel extraction kit (Qiagen), digested and used for restriction enzyme analysis by standard procedures (12).

Confirmation of Excision Foot Print by PCR Amplification and Sequencing. Template DNA from *niaD*436 was used in a PCR reaction in an attempt to amplify both the larger *niaD* sequence with an insert and the shorter *niaD* fragment resulting from excision of the Vader element. The PCR reaction was conducted as previously described, except for using primers MA003 (positions 359-378): 5'-

ATATGAATTCCTTCTTGACTTCCCCGGAAC-3' (SEQ ID NO:11) and NiaD5 (position 1125-1144): 5'-ATATAAGCTTGTCACTGGACGACATTTCAG-3' (SEQ ID NO:12). The gel purified fragment (ca. 800 bp) resulting from the excision event was submitted for sequencing.

Isolation of Tan1 via PCR Amplification. Fungal genomic DNA for PCR and Southern analyses was isolated from mycelia grown in CSL supplemented with 5% fructose (21). Genomic DNA of *A. niger* var. *awamori niaD* 436 mutant (22) was used as a template. A single primer (100 pmol), IR1, was used for amplification of Tan1. The 54-mer IR1 was derived from the 44 bp inverted repeat sequence of Vader preceded by a restriction enzyme recognition site for *EcoR*I: 5'-ATATGAATTC ACGTAATCAA CGGTCGGACG GGCCACACGG TCAGGCGGGC CATC-3' (SEQ ID NO:10). Deep Vent (exo⁻) DNA polymerase (New England Biolabs) was used with the buffer and dNTPs provided by the manufacturer. Denaturation of template DNA, 10 min. at 94°C, was followed by 30 cycles of denaturation (1 min. at 94°C), annealing of primers (1 min. at 55°C) and extension (6 min. at 72°C). PCR fragments were purified from agarose gels

using the Qiaex DNA gel extraction kit (Qiagen) and subcloned as blunt-ended inserts into *Eco*RV cut pSL1180 (Pharmacia Biotech).

Estimation of *niaD* Mutant Reversion Frequency. Spores from *niaD* mutants *niaD*392, *niaD*410, *niaD*436 and *niaD*587 were streaked onto minimal media containing NaNO₃ as a sole nitrogen source. Nitrate non-utilizing colonies of *niaD* mutants, which had a spidery appearance and did not sporulate, were streaked onto CM containing 600 mM potassium chlorate (KClO₃) and incubated to confluency at 37°C. Ten-fold dilution series of spore suspensions (in 0.8% NaCl-0.25% Tween 80) of *niaD*392, *niaD*410, *niaD*436, *niaD*587 and UVK143f wild-type spores were plated on minimal media with nitrate (10 mM) to determine reversion frequency, and on CM to determine viability.

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Southern Analysis. Genomic DNA for PCR and Southern analysis was isolated (13) from mycelia grown in CSL (13), which contained 600 mM KCIO₃ in order to reduce reversion of *niaD* back to the wild-type during cultivation. DNA (10 μg) was digested with either BgIII, which leaves the insertion intact in the *niaD* gene, or with EcoRV, which cuts the insertion element (Vader) once, and thus enables determination of its copy number in the genome. Genomic DNA (approximately 10 μg) of *A. nidulans, A. cinnamomeus, A. versicolor, A. wentii, A. phoenicis, A. foetidus* and of an industrial *A. niger* strain were digested with EcoRV to obtain an estimate of Vader copy number in these fungal genomes. The digested and gel-separated DNA was transferred to a positively-charged nylon membrane (Boehringer Mannheim) by capillary action.

The DNA probe for the *niaD* gene was derived from the PCR product (UVK143f DNA template amplified with primers NiaD1 (SEQ ID NO:8) and NiaD2 (SEQ ID NO:9)), which was digested with Sall, resulting in a 528 bp probe fragment. The probe for the insertion element, Vader, was derived from a PCR reaction in which *niaD436* DNA was used as a template. This PCR product was purified and digested with Sall and SphI and subcloned into the vector pUC19. This subclone was digested with Scal and Xbal to yield a 236 bp fragment which was used for estimation of the copy number of Vader sequences in the genomes of various fungi.

A DNA labeling and detection kit (Genius1, Boehringer Mannheim) was used for random primed labeling of probe DNA with digoxigenin, and for detection with alkaline-phosphatase labeled antibody to digoxigenin.

Hybridization and washing conditions for homologous probes were conducted as recommended by the manufacturer using hybridization buffer without formamide at 68°C (Boehringer Mannheim). Hybridizations for heterologous Southern analysis (i.e., analysis

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of DNA from other *Aspergillus* sp.) was conducted using hybridization buffer with 25% formamide at 37°C. Washes were performed as in stringent wash protocol.

Nitrate Reductase Assays. Nitrate reductase assays were performed as described in Dunn-Coleman, et al. (18).

DNA Analysis and Sequence Determination. Sequences were determined using fluorescent-labeled dideoxynucleotide terminators and *Taq* cycle sequencing on the 373A sequencer (ABI). Commercially available universal and reverse (New England Biolabs) primers were used. Alignment of sequences and prediction of amino acid sequences were performed using DNASTAR (DNASTAR, Inc.). The nucleotide and deduced amino acid sequences were analyzed and compared to those in GenBank, EMBL and Prot-Swiss using Fast A and BLAST programs (Genetics Computer Group, Inc. software package, Madison, WI).

Other Probes Used for Southern Analysis. The Tan1 probe was prepared by digesting Tan1 with *Hind*III and *Stu*I resulting in a 650 bp fragment corresponding to the 3' end of the transposase coding region (ORF-probe in Fig. 9). The Vader element was digested with *Xba*I and *Sca*I to yield a 236 bp fragment to be used for recognition of internal Vader sequence in Southern analysis (Vader-probe in Fig. 9).

Southern Analysis to Determine Tan1 Copy Number. Aspergillus genomic DNA (10 µg) was digested with *EcoRI*, which cuts the Tan1 element once in the transposase coding region and upstream of sequences corresponding to the Vader and Tan1 probes used in hybridizations (Figs. 5, 9 and 10). DNA labeling and detection kit (Genius 1, Boehringer Mannheim) was used for random primed labeling of probe DNA with digoxigenin and for detection with alkaline-phosphatase labeled antibody to digoxigenin. Hybridization and washing conditions were conducted as recommended by the manufacturer (Boehringer Mannheim).

Isolation of Tan1 from a Partial Genomic Library. It was known from the sequence of the PCR-amplified Tan1 element that Tan1 did not have restriction enzyme recognition sites for *Bgl*II and *Xho*I. A *Bgl*III-*Xho*I digested Southern blot of *Aspergillus niger* var. *awamori* genomic DNA, hybridized with the 650 bp *Hind*III-*Stu*I Tan1 probe, resulted in identification of a 4.5 kb genomic fragment containing Tan1. *A. niger* var. *awamori niaD*436 DNA was digested with *Bgl*II and *Xho*I and fragments in a size range of 4-5 kb were cloned into pSP73 (Promega). This partial genomic library was screened by colony hybridization using the nonradioactive nucleic acid labeling and detection system from Boehringer Mannheim.

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Example 1

Isolation of Spontaneous High Frequency Reverting niaD Mutants of A. niger var. awamon

Assuming that *niaD* mutants which arise from the insertion of a transposable element would be unstable, a total of 152 *niaD* mutants, isolated on the basis of spontaneous resistance to chlorate were characterized. To determine if the *niaD* mutation was unstable, spores from 43 *niaD* mutants were plated onto medium with nitrate as the sole nitrogen source. Fourteen of the mutants reverted to the wild-type phenotype at a frequency of greater than 1 X 10 ⁵. Table 1 summarizes the *niaD* mutant reversion studies.

Table 1

| | <u>Mutant</u> | Conidia Plated <u>No. x 10³</u> | No. Wild-Type <u>Colonies</u> | Reversion Frequency <u>x 10⁻⁴</u> |
|---|---------------|---|----------------------------------|--|
| i | niaD392 | 2.9 | 27 | 93 |
| | niaD410 | 7.7 | 5 | 6.5 |
| • | niaD436 | 3.7 | 164 | 443 |
| | niaD587 | 18.9 | 12 | 6.3 |

There appeared to be two classes of *niaD* mutants which reverted at high frequency. The *niaD* mutants *niaD*436 and *niaD*392 reverted at high frequency, while mutants *niaD*410 and *niaD*587 yielded smaller numbers of revertant colonies.

The level of nitrate reductase activity was determined using the assay described in (18) from revertant colonies isolated from the *niaD* 436 mutant. Nitrate reductase activity was detected in 14 of 15 revertants analyzed (see Table 2). A spectrum of activities was detected, suggesting that excision of Vader may not always be precise.

Table 2

| Strain | % Nitrate Reductase Activity Compared to Wild-Type |
|------------------------|--|
| UVK143f (wild-type) | 100 |
| niaD436 (niaD mutant) | ND ¹ |
| Revertants of niaD436: | |
| 1 | 34.7 |
| 2 | 42.8 |
| 3 | 27.7 |
| 4 | 3.5 |
| 5 | ND¹ |

¹ Activity non-detectable

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| ~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~ | *************************************** | % Nitrate Reductase Activity |
|---|---|------------------------------|
| | Strain | Compared to Wild-Type |
| 6 | | 47.4 |
| 7 | | 90.4 |
| 8 | | 9.8 |
| 9 | | 25.4 |
| 10 | | 28.9 |
| 11 | | 38.2 |
| 12 | | 6.9 |
| 13 | | 71.7 |
| 14 | | 71 .7 |
| 15 | | 49.7 |

Example 2 Cloning of a Vader Element

To determine if an insertion sequence was located within the *niaD* gene, two primers were synthesized. The first primer, niaD1 (SEQ ID NO:8), corresponded to position 142-165 of the *niaD* gene, and niaD2 (SEQ ID NO:9) corresponded to position 2738-2715 of the *niaD* gene. Genomic DNA was isolated from 14 unstable *niaD* mutants. This genomic DNA served as a template for the PCR primers. PCR reaction products with 4 *niaD* mutants (410, 436, 587 and 392) revealed an approximately 440 bp insertion (Vader) in the *niaD* gene.

For Southern blot analysis, genomic DNA isolated from the wild-type and four *niaD* mutants (410, 436, 587 and 392) was digested with Bglll. The probe used was a Sall digestion fragment of the 500 bp PCR product generated using the niaD1 (SEQ ID NO:8) and niaD2 (SEQ ID NO:9) oligomeric probes. The probe hybridized to a 2.5 kb fragment with wild-type DNA (lane 5, Fig. 1). In the case of the *niaD* mutants 410 (lane 1, Fig. 1), 436(lane 3, Fig. 1) and 392 (lane 4, Fig. 1), the probe hybridized to a 2.9 kb fragment. These results indicate that these three *niaD* mutants contain an approximately 440 bp insertion. Interestingly, with the mutant *niaD*587, the probe hybridized to both a 2.5 kb and 2.9 kb fragment, although mycelium had been grown in the experiment in the presence of KCIO₃ to favor growth of the *niaD* mutant and not revertant cells, the detection of two hybridizable sequences indicated that in some cells Vader had been excised from the *niaD* gene.

The approximate location of the insertion was determined in each of the four unstable *niaD* mutants by restriction mapping analysis. The location of the insertion in each of the four mutants examined is shown in Fig. 2. All four mutants had an approximately 440 bp insertion located at different sites within the *niaD* gene.

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Example 3

Determination of Vader Copy Number

To determine the Vader copy number a 236 bp Scal-Xbal internal fragment of Vader-2 (cloned from the mutant *niaD*436) was hybridized to EcoRV cleaved genomic DNA. There is only one EcoRV site within the Vader transposon. Southern blot analysis indicated that there are approximately fifteen copies of Vader sequences in the genome of *A. niger* var. *awamori*. (Fig. 4). The Vader sequences were integrated at identical genomic locations in the three *niaD* mutants, 410, 436 and 587. However, in the *niaD*392 mutant, Vader sequences were located in five different locations compared to the three *niaD* mutants examined. This result was somewhat surprising considering that all four *niaD* mutants were isolated from the same strain, but provides good evidence for the high mobility of the Vader element in this strain. When a propriety *A. niger* glucoamylase production strain (ETC #2663) was also examined, approximately 15 hybridization signals could be detected. Although some of the hybridization patterns appeared to be identical, clear differences could be seen between *A. niger* var. *awamori* and *A. niger*.

Example 4

Isolation of Vader in Other Fungal Species

In an attempt to determine if this transposable element was found in other filamentous fungi, genomic Southern blot analysis was performed using the 236 bp fragment (Xbal-Scal) of Vader sequence as per Example 3, as a probe (Fig. 5). Two strains of *A. nidulans* were obtained from Fungal Genetics Stock Center (FGSC), FGSC #A691, a nitrate reductase structural gene mutant (*niaD*15), and FGSC #A237, a tryptophan-requiring mutant (*trpC*801). No hybridization signals could be visualized with strain A691, and a single strong hybridization signal could be detected with strain A237. These results support the notion that the lack of success in cloning transposable elements from laboratory strains of *A. nidulans* is due to low copy number or absence. Similarly, only one hybridization signal could be detected in *A. foetidus* and *A. phoenicis*, while two hybridization signals were detected in *A. cinnamomeus*. No hybridizations could be detected in *A. wentii* and *A. versicolor*. In addition, no hybridization signals could be detected with *Humicola grisea* var. *thermoidea*, *Neurospora crassa* and *Trichoderma reesei* (results not shown). These results indicate that the Vader element is most commonly found in *A. niger* var. *awamori* and *A. niger*.

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Example 5

Excision of the Vader Element

Part of the *niaD* gene from *niaD*436 containing the Vader element was amplified using PCR. The PCR amplification resulted in the expected 1200 bp fragment of the Vader element flanked by *niaD* sequences and a shorter 800 bp fragment resulting from the excision event. Sequencing of the shorter fragment indicated that the Vader element had excised precisely. However, when several revertants of *niaD*436 and *niaD*410 were assayed for their nitrate reductase activity (18), a spectrum of activities was detected, suggesting that excision of the Vader element may not always be precise (results not shown).

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Example 6

Isolation of Tan1

The previously isolated Vader element, although mobile, did not have an ORF encoding transposase activity presumed to be required for excision (22). This observation led to a search for a transposase-encoding larger element, thus an oligomer corresponding to the Vader inverted repeat was synthesized and used for PCR amplification of the genomic *A. niger* var. *awamori* DNA. The PCR amplification resulted in the generation of three DNA fragments: the 0.4 kb Vader element, as expected, and fragments of 1.9 kb and 2.3 kb in length.

Both of the larger PCR-generated fragments were sequenced and the sequences were identical with an exception that the 2.3 kb fragment had an additional 400 bp at the 3' end. Surprisingly this additional sequence at the 3' end was a Vader element, which differed only by a few nucleotides from the previously isolated Vader. The 5' end sequence, shared by both of the 1.9 kb and 2.3 kb fragments, had a single ORF (1668 bp) coding for a protein of 555 amino acids flanked by inverted repeats (IRs). Thus, the 1.9 kb fragment, devoid of the Vader element, had an organization of IR-ORF-IR. The larger 2.3 kb fragment had a unique organization, IR-ORF-IR-Vader-IR, with a total of four inverted repeats (Figs. 9 and 11). In this larger element the two central inverted repeats, side by side, potentially form a tight hairpin structure, and despite many sequencing attempts with varying conditions, we were unable to determine the sequence between the two inverted repeats. However, the overall length of the PCR product, as determined by electrophoresis, corresponded to the size of the sequence shown in Fig. 11, suggesting that the two central contiguous IRs are not separated by a large segment of DNA.

Due to the organization of the 1.9 kb and 2.3 kb fragments, it was believed that the 1.9 kb fragment could have arisen in PCR from a partial amplification of the 2.3 kb

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fragment if the 3' IR-primer had annealed to the first central IR instead of the IR in the end of the Vader element. Southern analysis was conducted in order to determine if the 1.9 kb element existed in the genome without the associated Vader element, or whether it was a PCR-artifact derived from a partial amplification of the 2.3 kb element. The two probes used in Southern analysis corresponded to the internal sequence of Vader and to the carboxyterminal part of the ORF (Fig. 9). The genomic DNA from *A. niger* var. awamon niaD mutants and UVK143f were digested with EcoRI, which cuts once in the coding region of the ORF upstream from the ORF-probe and does not cut Vader. The Southern analysis showed numerous bands for the Vader element (Fig. 10), similar to previous Southern analyses (22). However, only one fragment lit up with the probe corresponding to the ORF and a fragment of the same size (1.6 kb) was recognized by the Vader probe (Fig. 10). It was concluded that the actual element in the genome was the 2.3 kb fragment and that the shorter 1.9 kb had only been a PCR-artifact. The isolated 2.3 kb fragment was designated as Tan1.

A genomic clone of the Tan1 element (2.3 kb) was isolated from a partial genomic library. Restriction enzymes, which were shown not to have any recognition sites in the PCR-amplified Tan1, were used separately and in combinations in Southern analysis of the genomic DNA. A double digestion with *Bg/*III and *Xho*I resulted in a relatively short, 4.5 kb, fragment which hybridized with the ORF-specific probe (data not shown). Genomic DNA fragments cleaved by *Bg/*III and *Xho*I and between 4 kb and 5 kb in size were cloned into pSP73 (Promega). The correct clone containing the Tan1 element was isolated by colony hybridization using the ORF-specific probe. Differences between the sequences of the genomic clone and the PCR-generated Tan1 were minor, even for the flanking IRs which were almost identical even though in the PCR-generated Tan1 the IRs were derived from the Vader IRs (PCR primers). It was seen from the genomic clone of Tan1 that immediately outside of the terminal IRs there were TA-dinucleotides, suggesting a TA target site and its duplication upon insertion. Sequence of the Tan1 genomic clone is shown in Figs. 11A and 11B [SEQ ID NO:13 (DNA) and SEQ ID NO:14 (amino acid)].

Example 7

Insertional Inactivation/Gene Tagging

Vader was cloned by insertional inactivation of the target gene *niaD*, which encodes nitrate reductase. The target sequence for integration of Vader is TA, a sequence which must be very common in the genome of fungi. Nitrate reductase mutants cannot grow on nitrate and inconsequence are resistant to the toxic analog of nitrate, KCIO₃.

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It is possible that one of the reasons heterologous protein production in fungi is lower than that of homologously produced protein using the same promoter is that the heterologous protein is being degraded by the cell. If there are genes whose products are responsible for degrading/sequestering foreign protein, it would be advantageous to inactivate those genes. In order to achieve this, a strain is constructed using gene disruption, which lacks the Tan1 gene. Such strain is then used to transform and express a heterologous protein such as the mammalian chymosin protein. It would be advantageous if the activity of such genes could be visualized or selected for on petri dishes. For example chymosin produced in *A. niger* results in a halo of clearing around a colony grown on skim milk. (See US Patent 5,364,770, the disclosure of which is incorporated herein by reference.)

Having transformed the strain with a construct comprising the desired heterologous protein or polypeptide, one would transform the strain a second time with Vader and Tan1 appropriately modified for gene tagging purposes.

The transformants are then plated on medium which can be used to visualize heterologous protein production, such as skim milk plates in the case of chymosin.

The plates are then screened for increased halo size, which is the result of inactivation of a gene whose product limits foreign protein production.

The inactivated gene can be cloned using the transposon sequences as a marker for cloning strategies. (See generally (19),)

Example 8

Elevation of Gene Expression Using Transposons

A reason that heterologous protein production is lower than expected in fungi is presumed to be that genes essential for foreign (heterologous) gene production are NOT expressed at sufficiently high levels in the fungi.

In order to overcome this problem, utilizing the transposable element(s) of the present invention, a strain is constructed in which the native Tan1 gene is inactivated by gene disruption.

This strain is used to express a heterologous protein whose expression can be easily visualized, such as chymosin (US Patent 5,364,770). A second transformation is made with Vader and Tan1, appropriately modified for gene tagging purposes. The internal sequence of Vader is replaced by a promoter sequence. One of the many integration events possible will be the integration of this promoter carrying Vader element into 5' to a gene beneficial to heterologous protein (e.g., chymosin) expression or secretion. Upon insertion, this beneficial gene is activated and such integrant colonies can be screened for,

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e.g., increased halo size (chymosin). The activated gene can be cloned using the transposon sequences as a marker for cloning strategies.

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Sequence Listing

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: Amutan, Maria

Dunn-Coleman, Nigel Nyyssonen, Eini M.

- (ii) TITLE OF INVENTION: Identification of and Cloning a Mobile Transposon from Aspergillus
- (iii) NUMBER OF SEQUENCES: 17
- (iv) CORRESPONDENCE ADDRESS:
 - (A) NAME: Genencor International, Inc.
 - (B) STREET: 925 Page Mill Road
 - (C) CITY: Palo Alto
 - (D) STATE: CA
 - (E) COUNTRY: USA
 - (F) POSTAL CODE (ZIP): 94304
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.25 (EPO)
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US
 - (B) FILING DATE: August 16, 1996
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Horn, Margaret A.
 - (B) REGISTRATION NUMBER: 33,401
 - (C) REFERENCE/DOCKET NUMBER: GC270-2
 - (xi) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (415) 846-7536
 - (B) TELEFAX: (415) 845-6504
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 45 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

ACGTAATCAA CGGTCGGGCG GGCCACACGG TCAGGCGGGC CACCC

(2) INFORMATION FOR SEQ ID NO: 2:

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(ii) MOLECULE TYPE: DNA

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| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: DNA | |
|---|-----|
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2: | |
| GATGGCCCGC CTGACCGTGT GGCCCGCCCG ACCGTTGATT ACGT | 44 |
| (2) INFORMATION FOR SEQ ID NO: 3: | |
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 437 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |
| (ii) MOLECULE TYPE: DNA | |
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3: | |
| ACGTAATCAA CGGTCGAACG GGCCACACGG TCAGGCGGGC CATCCTGAAA TCCCATATAA | 60 |
| AAGATGTCTT GGGGATTCTA TTATATATCA ACCAGTACTA CTTCTATGAA GCTCTAACTT | 120 |
| TGTAGATAGT TATATATA AGAATAAGTA TTCCATGAAT TTTTCAGATT TTAGAATTTT | 180 |
| TACTTTGATA ATGAAACCAG ATTCTTATAT AAAACATATA AATACAGATA TTGTAATATG | 240 |
| ATAAGTCCAT AAGTAAAAGT ATATTCATTT TTAGAAGGTA TATAGATATT ATTTATATTA | 300 |
| TTTAAAATCT ATATAGAAGA AATCTAATTC TTCTAGACCT GGATGGTAGA GATATATTAT | 360 |
| GTTTAAAAAG ATATCTTTTG TATAGTATTA CCAGATGGCC CGCCTGACCG TGTGGCCCGT | 420 |
| CCGACCGTTG ATTACGT | 437 |
| (2) INFORMATION FOR SEQ ID NO: 4: | |
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |
| (ii) MOLECULE TYPE: DNA | |
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4: | |
| ACGTAATCAA CGGTCGAACG GGCCACACGG TCAGGCGGGC CATC | 44 |
| (2) INFORMATION FOR SEQ ID NO: 5: | |
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 44 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

GATGGCCCGC CTGACCGTGT GGCCCGTCCG ACCGTTGATT ACGT

44

(2) INFORMATION FOR SEQ ID NO: 6:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2325 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

| 60 | AAAACACCAC | CATCCCTTCG | TCAGGCGGGC | GGCCACACGG | CGGTCGGACG | ACGTAATCAA |
|------|------------|------------|------------|------------|------------|------------|
| 120 | CATCTATCCC | CCACCAAAAG | ACCACAAATG | CTTTTCAACC | CTACCCGAGG | CTTGAATCAC |
| 180 | CTATTCAGAA | GCCATTGAAG | GATTCTTCTT | AGGAAGGCAG | CAGGTGGAGC | ATCAAAATCG |
| 240 | GAACTACTCT | GACGTCGCTC | GCGTGTTTAT | GTGAAGCAGC | ACTAGTATTC | AGGCCAAATC |
| 300 | GTCAAAAATT | ACCAACGCAC | TAAAAATATG | GTGTTTTCGC | TTATCTGGAC | CCAGGCTCGA |
| 360 | AGCGAGGAGC | TCTCTAGATA | ATGGATCCTA | CGCTTGTTAA | GAAGAGGAAT | GTCAAATAAT |
| 420 | AACGAGGTTA | ATTATCTCTA | GGCTAATTTG | TCAGAGATAT | CCACTTGATA | AAGCCCCCGG |
| 480 | GCCACGAATC | TTTGTTAAAC | GGCTTATAGC | GCATCAACTG | GAACAAGTAG | TTCAACTGTT |
| 540 | AGGATCCTGA | GCTAAAATGG | CTATCCAAGA | GACGACTCAA | CGATTTGCTA | CCTACGAACT |
| 600 | GGATCTCATC | CAAGAGTACG | GGAAGTTATT | AACGCGTACA | GACTGGTTCC | AGTTATAAAA |
| 660 | CTACATATAA | GGAATGATTG | GTTTGCTATG | ATGAAACAGG | TACAATTTCG | AGATGATATA |
| 720 | GGAATCGGGA | GTTCAACCAG | GCCGTCCCTA | GGGCAGGTCG | AGTTCCCAGA | AGTAGTAACT |
| 780 | CGACCCTGAT | GTTCTACCTT | TAATGGAGAG | GTATTCGCTC | CCAATTGAGT | ATGGGTCACT |
| 840 | CTCCTACCTG | CAATCTATTC | GTATGAAGGT | TAAAGGCATG | AAAACACATC | CTTTAAAGGC |
| 900 | GATGGCTTCC | ATTGGACTTC | TACTGATAAA | ATGGTTGGAC | GTCAGTGATA | GAGATTTGAA |
| 960 | TCCTAGTCCT | AAATATAGCC | ATCAGTAGGC | TTAGAGGCAA | ATTCCCTTGA | AAAACACTTC |
| 1020 | AAAATGAGGT | TCCTGTGCTG | ATTCGACCAA | TGACACCTGA | GGTAGTCATT | CGATGGCCAC |
| 1080 | ATGTTGGTTG | CAGCCTCTTG | CCATCTACTT | CTCATTCGTC | TGTATGCCAG | TATACCTATT |
| 1140 | AATACGGCCG | AAGCAGATGC | CATGGTTCCC | CGTACGGAGG | CTTAAACGCA | TTTTAGTGTG |
| 1200 | AGTGTGCTTT | AAAGCTCACC | GGTCTATCCT | ACTTCTTAGA | GACAAGCTTG | CAATCATATC |
| 1260 | TAGATCCTGA | CTTGTTCCTC | AGCAACAGGT | GTGGTTTTAG | AATATAATCA | ATCAAAGTCG |
| 1320 | ATAGCCAGTC | CCAACCCCGG | GAAAACACCA | ATATTCGCTT | TCTCGACTCC | TCAAGTGCTT |
| 1380 | ATCCAAAATC | CTTTTGGAGC | TATAAAACAC | CACCACATAA | GTGCTTCAAA | AAGTGGCTCA |
| 1440 | ACTCTACACT | TCACCTACAA | AAGTCCAACT | AACGGCAAGC | CTACTTCGGA | AGTGGAACGC |
| | | | | | | |

| ACGTCAGCTT | CTCAAAGGGT | GTGAACTAGC | AATAACAAAC | TCAATCATAC | TGGCTAAGGA | 1500 |
|------------|------------|------------|------------|------------|------------|------|
| GAATGCGGAA | TTACGTGCTA | GCCATGAAAA | GCAACTACCA | AAGAGGAAGC | GTTCAAGGAA | 1560 |
| GCAGGTGATC | TATACAGAAG | GCACTACCGT | TGAAGAGGCC | CAGAGAGCTA | TACAGGAAGT | 1620 |
| GGAAGAGGTG | CAGAATGATG | AAGATATTGA | GGTTGAACCC | CAATCTCAAT | ATACGGAGAC | 1680 |
| CCCCTCGCGC | GCGCCTCCAC | GCTGCAGTAA | TTGCTTCAAT | ATAGGCCACC | GACGTACACA | 1740 |
| GTGTTCTAAA | CCACCTACTA | ATTAGTTAGA | TAGCTGTTTT | TACAAGCATT | TATGTTGATT | 1800 |
| TAGAGGCCTC | ATTTGGATCA | TATCGGGTAA | TCCTACCGGG | AGATGGCCCG | CCTGACCGTG | 1860 |
| TGGCCCGCCC | GACCGTTGAT | TACGTNNNNN | ACGTAATCAA | CGGTCGGACG | GGCCCCCGG | 1920 |
| TCCGGCGGGC | CATCTGGTAA | TACTATACCA | AAGATATCTT | TTTAAACATA | ATATATCTCT | 1980 |
| ACCATCCAGG | TCTAGGAGAA | TTAGATTTCT | TCTATATAGA | TTTTAAATAA | TATAAATAAT | 2040 |
| ATCTATATAC | CTTCTAAAAA | TGAATATACT | TTTACTTATG | GACTTATCAT | ATTACAATAT | 2100 |
| CTGTATTTAT | ATGTATTATA | TAAGAATCTG | GTTTCATTAT | CAAAGTAAAA | ATTCTAAAAT | 2160 |
| CTGAAAAATT | CATGGAATAC | TTATTCTTAT | ATATATAACT | ATCTACAAAG | TTAGAGCTTC | 2220 |
| ATAGAAGTAG | TACTGGTTGA | TATATAATAG | AATCCCCAAG | ACATCTTTTA | TATGGGATTT | 2280 |
| CAGGATGGCC | GCCGACCGTG | TGGCCCGTCC | GACCGTTGAT | TACGT | | 2325 |

(2) INFORMATION FOR SEQ ID NO: 7:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 555 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:

Met Pro Pro Lys Ala Ser Ile Pro Ser Lys Ser Gln Val Glu Glu 1 5 10 15

Gly Arg Ile Leu Leu Ala Ile Glu Ala Ile Gln Lys Gly Gln Ile Thr 20 25 30

Ser Ile Arg Glu Ala Ala Arg Val Tyr Asp Val Ala Arg Thr Thr Leu
35 40 45

Gln Ala Arg Leu Ser Gly Arg Val Phe Ala Lys Asn Met Thr Asn Ala 50 55 60

Arg Gln Lys Leu Ser Asn Asn Glu Glu Glu Ser Leu Val Lys Trp Ile 65 70 75 80

Leu Ser Leu Asp Lys Arg Gly Ala Ser Pro Arg Pro Leu Asp Ile Arg 85 90 95

Asp Met Ala Asn Leu Ile Ile Ser Lys Arg Gly Tyr Ser Thr Val Glu 100 105 110

| Gln | Val | Gly 115 | Ile | Asn | Trp | Ala | Туг 120 | Ser | Phe | Val | Lys | Arg 125 | | Glu | Ser |
|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| Leu | Arg 130 | Thr | Arg | Phe | Ala | Arg 135 | Arg | Leu | Asn | Tyr | Pro 140 | Arg | Ala | Lys | Met |
| Glu 145 | Asp | Pro | Glu | Val | Ile 150 | Lys | Asp | Trp | Phe | Gln 155 | Arg | Val | Gln | Glu | Val 160 |
| Ile | Gln | Glu | Tyr | Gly 165 | Ile | Ser | Ser | Asp | Asp 170 | Ile | Tyr | Asn | Phe | Asp 175 | Glu |
| Thr | Gly | Phe | Ala 180 | Met | Gly | Met | Ile | Ala 185 | Thr | Tyr | Lys | Val | Val 190 | Thr | Ser |
| Ser | Gln | Arg 195 | Ala | Gly | Arg | Pro | Ser 200 | Leu | Val | Gln | Pro | Gly 205 | Asn | Arg | Glu |
| Trp | Val 210 | Thr | Pro | Ile | Glu | Cys 215 | Ile | Arg | Ser | Asn | Gly 220 | Glu | Val | Leu | Pro |
| Ser 225 | Thr | Leu | Ile | Phe | Lys 230 | Gly | Lys | Thr | His | Leu 235 | Lys | Ala | Trp | Tyr | Glu 240 |
| Gly | Gln | Ser | Ile | Pro 245 | Pro | Thr | Trp | Arg | Phe 250 | Glu | Val | Ser | Asp | Asn 255 | Gly |
| Trp | Thr | Thr | Asp 260 | Lys | Ile | Gly | Leu | Arg 265 | Trp | Leu | Pro | Lys | His 270 | Phe | Ile |
| Pro | Leu | 11e 275 | Arg | Gly | Lys | Ser | Val 280 | Gly | Lys | Tyr | Ser | Leu 285 | Leu | Val | Leu |
| Asp | Gly 290 | His | Gly | Ser | His | Leu 295 | Thr | Pro | Glu | Phe | Asp 300 | Gln | Ser | Cys | Ala |
| Glu 305 | Asn | Glu | Val | Ile | Pro 310 | ļlе | Cys | Met | Pro | Ala 315 | His | Ser | Ser | His | Leu 320 |
| Leu | Gln | Pro | Leu | Asp 325 | Val | Gly | Cys | Phe | Ser 330 | Val | Leu | Lys | Arg | Thr 335 | Tyr |
| Gly | Gly | Met | Val 340 | Pro | Lys | Gln | Met | Gln 345 | Tyr | Gly | Arg | Asn | His 350 | Ile | Asp |
| Lys | Leu | Asp 355 | Phe | Leu | Glu | Val | Tyr 360 | Pro | Lys | Ala | His | Gln 365 | Cys | Ala | Leu |
| Ser | Lys 370 | Ser | Asn | Ile | Ile | Ser 375 | Gly | Phe | Arg | Ala | Thr 380 | Gly | Leu | Val | Pro |
| Leu 385 | Asp | Pro | Asp | Gln | Val 390 | Leu | Ser | Arg | Leu | His 395 | Ile | Arg | Leu | Lys | Thr 400 |
| Pro | Pro | Thr | Pro | Asp 405 | Ser | Gln | Ser | Ser | Gly 410 | Ser | Val | Leu | Gln | Thr 415 | Pro |
| His | Asn | Ile | Lys 420 | His | Leu | Leu | Glu | His 425 | Pro | Lys | Ser | Val | Glu 430 | Arg | Leu |
| Leu | Arg | Lys 435 | Arg | Gln | Ala | Ser | Pro | Thr | Ser | Pro | Thr | Asn 445 | Ser | Thr | Leu |

| | Arg | 450 | Leu | Leu | гÀг | сту | 455 | GIU | Leu | MIG | 116 | 460 | ASII | JCI | 110 | | |
|-------|------------|------------|-------------------------------|-----------------------|----------------------|-------------------|----------------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|----|
| | Leu 465 | Ala | Lys | Glu | Asn | Ala 470 | Glu | Leu | Arg | Ala | Ser 475 | His | Glu | Lys | Gln | Leu 480 | |
| | Pro | Lys | Arg | Lys | Arg 485 | Ser | Arg | Lys | Gln | Val 490 | Ile | Tyr | Thr | Glu | Gly 495 | Thr | |
| | Thr | Val | Glu | Glu 500 | Ala | Gln | Arg | Ala | Ile 505 | Gln | Glu | Val | Glu | Glu 510 | Val | Gln | |
| | Asn | Asp | Glu 515 | Asp | Ile | Glu | Val | Glu 520 | Pro | Gln | Ser | Gln | Tyr 525 | Thr | Glu | Thr | |
| | Pro | Ser 530 | Arg | Ala | Pro | Pro | Arg 535 | Cys | Ser | Asn | Суѕ | Phe 540 | Asn | Ile | Gly | His | |
| | Arg 545 | Arg | Thr | Gln | Cys | Ser 550 | Lys | Pro | Pro | Thr | Asn 555 | | | | | | |
| (2) | INFO | RMAT: | ION : | FOR : | SEQ : | ID N | o: 8 | : | | | | | | | | | |
| | (i) | (B (C | UENC:) LEI) TY:) ST:) TO: | NGTH PE: 1 RAND | : 23 nucl EDNE | bas eic SS: | e pa acid sing | irs | | | | | | | | | |
| | (ii) | MOL | ECUL: | E TY | PE: | AND | | | | | | | | | | | |
| | (xi) | SEQ | UENC | E DE | SCRI | PTIO | N: S | EQ I | D NO | : 8: | | | | | | | |
| CCA | ACCGA | GT C | CTCA | GTAT. | A GA | С | | | | | | | | | | | 23 |
| (2) | INFO | RMAT | ION | FOR | SEQ | ID N | 0: 9 | : | | | | | | | | | |
| | (i) | (B (C | UENC) LE) TY) ST) TO | NGTH PE: RAND | : 24 nucl EDNE | bas eic SS: | e pa acid sing | irs | | | | | | | | | |
| | (ii) | MOL | ECUL | E TY | PE: | DNA | | | | | | | | | | | |
| | (xi) | SEQ | UENC | E DE | SCRI | PTIO | N: S | EQ I | D NO | : 9: | | | | | | | |
| CAA | GCTT | CA T. | AGGC | GTCC. | A GA | TC | | | | | | | | | | | 24 |
| (2) | INFO | RMAT | ION | FOR | SEQ | ID N | 0: 1 | 0: | | | | | | | | | |
| | (i) | (B (C | UENC) LE) TY) ST) TO | NGTH PE: RAND | : 54 nucl EDNE | bas eic SS: | e pa acid sing | irs | | | | | | | | | |
| | (ii) | MOL | ECUL | E TY | PE: | DNA | | | | | | | | | | | |
| | (xi) | SEQ | UENC | E DE | SCRI | PTIO | N: S | EQ I | D NO | : 10 | : | | | | | | |
| ימידמ | rgaat | TC A | .CGTA | ATCA | A CG | GTCG | GACG | GGC | CACA | CGG | TCAG | GCGG | GC C | ATC | | | 54 |

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| (2) INFORMATION FOR SEQ ID NO: 11: | |
|--|-----|
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |
| (ii) MOLECULE TYPE: DNA | |
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11: | |
| ATATGAATTC CTTCTTGACT TCCCCGGAAC 30 | |
| (2) INFORMATION FOR SEQ ID NO: 12: | |
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |
| (ii) MOLECULE TYPE: DNA | |
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12: | |
| ATATAAGCTT GTCACTGGAC GACATTTCAG 30 | |
| (2) INFORMATION FOR SEQ ID NO: 13: | |
| (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2329 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear | |
| (ii) MOLECULE TYPE: DNA (genomic) | |
| (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13: | |
| ACGTAATCAA CGGTCGGGCG GGCCACACGG TCAGGCGGGC CACCCCTTCG AAAACACCAC | 60 |
| CTTGAATCAC CTACCCGAGG CTTTTCAACC ACCACAAATG CCACCAAAAG CATCTATCCC | 120 |
| ATCAAAATCG CAGGTGGAGC GGGAAGGCAG GATTCTTCTT GCCATTGAAG CTATTGAGAA | 180 |
| AGGCCAAATC ACTAGTATTC GTGAAGCAGC GCGTGTTTAT GACGTCGCTC GAACTACTCT | 240 |
| CCAGGCTCGA TTATCTGGAC GTGTTTTCGC TAAAAATATG ACCAACGCAC GTCAAAAATT | 300 |
| GTCAAATAAT GAAGAGGAAT CGCTTGTTAA ATGGATCCTA TCTCTAGATA AGCGAGGAGC | 360 |
| AAGCCCCCGG CCACTTGATA TCAGAGATAT GGCTAATTTG ATTATCTCTA AACGAGGTTA | 420 |
| TTCAACTGTT GAACAAGTAG GCATCAACTG GGCTTATAGC TTTGTTAAAC GCCACGAATC | 480 |
| CCTACGAACT CGATTTGCTA GACGACTCAA CTATCAAAGA GCTAAAATGG AGGATCCTGA | 540 |
| AGTTATAAAA GACTGGTTCA AACGCGTACA GGAAGTTATT CAAGAGTACG GGATCTCATC | 600 |
| AGATGATATA TACAATTTCG ATGAAACAGG GTTTGCTATG GGAATGATTG CTACATATAA | 660 |

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| AGTAGTAACT | AGTTCCCAGA | GGGCAGGTCG | GCCGTCCCTA | GTTCAACCAG | GGAATCGGGA | 720 |
|------------|------------|------------|------------|------------|------------|------|
| ATGGGTCACT | GCAATTGAGT | GTATTCGCTC | TAATGGAGAG | GTTCTACCTT | CGACCCTGAT | 780 |
| CTTTAAAGGC | AAAACACATC | TAAAGGCATG | GTATGAAGGT | CAATCTATTC | CTCCTACCTG | 840 |
| GAGATTTGAA | GTCAGTGATA | ATGGTTGGAC | TACTGATAAA | ATTGGACTTC | GATGGCTTCA | 900 |
| AAAACACTTC | ATTCCCTTGA | TTAGAGGCAA | ATCAGTAGGC | AAATATAGCC | TCCTAGTCCT | 960 |
| CGATGGCCAC | GGTAGTCATT | TGACACCTGA | ATTCGACCAA | TCCTGTGCTG | AAAATGAGGT | 1020 |
| TATACCTATT | TGTATGCCTG | CTCATTCGTC | CCATCTACTT | CAGCCTCTTG | ATGTTGGTTG | 1080 |
| TTTTAGTGTG | CTTAAACGCA | CGTACGGAGG | CATGGTTCAA | AAGCAGATGC | AATACGGCCG | 1140 |
| CAATCATATC | GACAAGCTTG | ACTTCTTAGA | GGTCTATCCT | AAAGCTCACC | AGTGTGCTTT | 1200 |
| ATCAAAGTCG | AATATAATCA | GTGGTTTTAG | AGCAACAGGT | CTTGTTCCTC | TAGATCCTGA | 1260 |
| TCAAGTGCTT | TCTCGACTCC | ATATTCGCTT | GAAAACACCA | CCAACCCCGG | ATAGCCAGTC | 1320 |
| AAGTGGCTCA | GTGCTTCAAA | CACCACATAA | TATAAAACAC | CTTTTGAAGC | ATCCAAAATC | 1380 |
| AGTGGAACGC | CTACTTCGGA | AACGGCAAGC | AAGTCCAACT | TCACCTACAA | ACTCTACACT | 1440 |
| ACGTCAGCTT | CTCAAAGGGT | GTGAACTAGC | AATAACAAAC | TCAATCATAC | TGGCTAAGGA | 1500 |
| GAATGCGGAA | TTACGTGCTA | GCCATGAAAA | GCAACTACCA | AAGAGGAAGC | GTTCAAGGAA | 1560 |
| GCAGGTGATC | TATACAGAAG | GCACTACCGT | TGAAGAGGCC | CAGAGAGCTA | TACAGGAAGT | 1620 |
| GGAAGAGGTG | CAGAATGATG | AAGATATTGA | GGTTGAACCC | CAATCTCAAT | ATACGGAGAC | 1680 |
| CCCCTCGCGC | GCGCCTCCAC | GCTGCAGTAA | TTGCTTCAAT | ATAGGCCACC | GACGTACACA | 1740 |
| GTGTTCTAAA | CCACCTACTA | ATTAGTTAGA | TAGCTGTTTT | TACAAGCATT | TATGTTGATT | 1800 |
| TAGAGGCCTC | ATTTTGATCA | TATCGGGTAA | TCCTACCGAG | AGATGGCCCG | CCTGACCGTG | 1860 |
| TGGCCCGCCC | GACCGTTGAT | TACGTNNNNN | ACGTAATCAA | CGGTCGGACG | GCCCCCCGG | 1920 |
| TCCGGCGGGC | CATCTGGTAA | TACTATACAA | AAGATATCTT | TTTAAACATA | ATATATCTCT | 1980 |
| ACCATCCAGG | TCTAGGAGAA | TTAGATTTCT | TCTATATAGA | TTTTAAATAA | TATAAATAAT | 2040 |
| ATCTATATAC | CTTCTAAAAA | TGAATATACT | TTTACTTATG | GACTTATCAT | ATTACAATAT | 2100 |
| CTGTATTTAT | ATGTATTATA | TAAGAATCTG | GTTTCATTAT | CAAAGTAAAA | ATTCTAAAAA | 2160 |
| TCTGAAAAAT | TCATGGAATA | CTTATTCTTA | ТАТАТАТАА | CTATCTACAA | AGTTAGAGCT | 2220 |
| TCATAGAAGT | AGTACTGGTT | GATATATAAT | AGAATCAAAA | AGACATCTTT | TATATGGGAT | 2280 |
| TTCAGGATGG | CCCGCCTGAC | CGTGTGGCCC | GTTCGACCGT | TGATTACGT | | 2329 |
| | | | | | | |

(2) INFORMATION FOR SEQ ID NO: 14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 555 amino acids
 (B) TYPE: amino acid
 (C) STRANDEDNESS: single

- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:
- Met Pro Pro Lys Ala Ser Ile Pro Ser Lys Ser Gln Val Glu Arg Glu 1 5 10 15
- Gly Arg Ile Leu Leu Ala Ile Glu Ala Ile Arg Lys Gly Gln Ile Thr 20 25 30
- Ser Ile Arg Glu Ala Ala Arg Val Tyr Asp Val Ala Arg Thr Thr Leu 35 40 45
- Gln Ala Arg Leu Ser Gly Arg Val Phe Ala Lys Asn Met Thr Asn Ala 50 60
- Arg Gln Lys Leu Ser Asn Asn Glu Glu Glu Ser Leu Val Lys Trp Ile 65 70 75 80
- Leu Ser Leu Asp Lys Arg Gly Ala Ser Pro Arg Pro Leu Asp Ile Arg 85 90 95
- Asp Met Ala Asn Leu Ile Ile Ser Lys Arg Gly Tyr Ser Thr Val Glu 100 105 110
- Gln Val Gly Ile Asn Trp Ala Tyr Ser Phe Val Lys Arg His Glu Ser 115 120 125
- Leu Arg Thr Arg Phe Ala Arg Arg Leu Asn Tyr Gln Arg Ala Lys Met 130 135 140
- Glu Asp Pro Glu Val Ile Lys Asp Trp Phe Lys Arg Val Gln Glu Val 145 150 155 160
- Ile Gln Glu Tyr Gly Ile Ser Ser Asp Asp Ile Tyr Asn Phe Asp Glu 165 170 175
- Thr Gly Phe Ala Met Gly Met Ile Ala Thr Tyr Lys Val Val Thr Ser 180 185 190
- Ser Gln Arg Ala Gly Arg Pro Ser Leu Val Gln Pro Gly Asn Arg Glu 195 200 205
- Trp Val Thr Ala Ile Glu Cys Ile Arg Ser Asn Gly Glu Val Leu Pro 210 215 220
- Ser Thr Leu Ile Phe Lys Gly Lys Thr His Leu Lys Ala Trp Tyr Glu 225 230 235 240
- Gly Gln Ser Ile Pro Pro Thr Trp Arg Phe Glu Val Ser Asp Asn Gly
 245 250 255
- Trp Thr Thr Asp Lys Ile Gly Leu Arg Trp Leu Gln Lys His Phe Ile 260 265 270
- Pro Leu Ile Arg Gly Lys Ser Val Gly Lys Tyr Ser Leu Leu Val Leu 275 280 285
- Asp Gly His Gly Ser His Leu Thr Pro Glu Phe Asp Gln Ser Cys Ala 290 295 300

Glu Asn Glu Val Ile Pro Ile Cys Met Pro Ala His Ser Ser His Leu Leu Gln Pro Leu Asp Val Gly Cys Phe Ser Val Leu Lys Arg Thr Tyr Gly Gly Met Val Gln Lys Gln Met Gln Tyr Gly Arg Asn His Ile Asp Lys Leu Asp Phe Leu Glu Val Tyr Pro Lys Ala His Gln Cys Ala Leu Ser Lys Ser Asn Ile Ile Ser Gly Phe Arg Ala Thr Gly Leu Val Pro Leu Asp Pro Asp Gln Val Leu Ser Arg Leu His Ile Arg Leu Lys Thr Pro Pro Thr Pro Asp Ser Gln Ser Ser Gly Ser Val Leu Gln Thr Pro 410 His Asn Ile Lys His Leu Leu Lys His Pro Lys Ser Val Glu Arg Leu Leu Arg Lys Arg Gln Ala Ser Pro Thr Ser Pro Thr Asn Ser Thr Leu Arg Gln Leu Leu Lys Gly Cys Glu Leu Ala Ile Thr Asn Ser Ile Ile Leu Ala Lys Glu Asn Ala Glu Leu Arg Ala Ser His Glu Lys Gln Leu Pro Lys Arg Lys Arg Ser Arg Lys Gln Val Ile Tyr Thr Glu Gly Thr Thr Val Glu Glu Ala Gln Arg Ala Ile Gln Glu Val Glu Val Gln Asn Asp Glu Asp Ile Glu Val Glu Pro Gln Ser Gln Tyr Thr Glu Thr Pro Ser Arg Ala Pro Pro Arg Cys Ser Asn Cys Phe Asn Ile Gly His 535 Arg Arg Thr Gln Cys Ser Lys Pro Pro Thr Asn

(2) INFORMATION FOR SEQ ID NO: 15:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 44 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

ACGTAATCAA CGGTCGGACG GGCCCCCCGG TCAGGCGGGC CATC

(2) INFORMATION FOR SEQ ID NO: 16:

| (i) | SEQUENCE | CHARAC' | TERISTI | CS |
|-----|------------|---------|---------|----|
| | (B) T 7717 | | - | _ |

- (A) LENGTH: 45 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

GGATGGCCCG CCTGACCGTG TGGCCCGTTC GACCGTTGAT TACGT

45

- (2) INFORMATION FOR SEQ ID NO: 17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 44 base pairs

 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

ACGTAATCGG TAAGCGAGTT GCCCGCGCAA GCGAGTTGCC CACC

44

What is Claimed

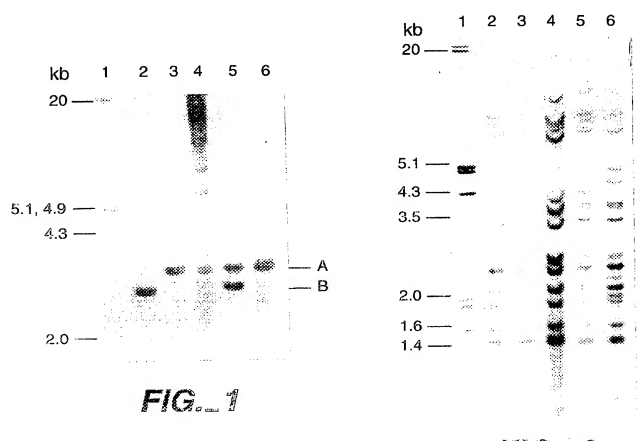
- 1. A transposable element isolated from *Aspergillus niger* var. awamori comprising a DNA fragment of about 2.3 kb which comprises SEQ ID NO: 1.
- The transposable element of Claim 1 comprising the DNA sequence of SEQ ID NO:13 or variations thereof.
- 3. A fragment of the transposable element of Claim 1 comprising part or all of the DNA sequence selected from the group consisting of SEQ ID NOS:1 and 16 or variations thereof.
- 4. An isolated transposase coded for by the transposable element of Claim 1.
- 5. The transposase of Claim 4 comprising the amino acid sequence of SEQ ID NO:14.
- 6. A method of isolating a transposable element from a filamentous fungus, comprising the steps of:
 - (a) hybridizing fungal DNA under low stringency conditions to a probe, wherein the probe comprises part or all of one of the DNA fragments of Claim 3; and
 - (b) isolating fungal DNA which hybridizes to said probe.
- 7. The method of Claim 6 wherein the probe comprises an imperfect direct repeat within the DNA sequence selected from the group consisting of SEQ ID NOS:1 and 16.
- 8. A method of isolating a transposable element from a filamentous fungus genomic library, the method comprising probing said library with an ORF-specific probe and isolating DNA which hybridizes to said ORF-specific probe.
- A method of isolating a transposable element from a filamentous fungi, the method comprising,
 - (a) subjecting fungal DNA to polymerase chain reaction amplification using part or all of one of the DNA fragments of Claim 3 as a primer, thereby generating amplified DNA sequences;
 - (b) isolating the amplified DNA sequences; and
 - (c) optionally identifying said amplified DNA sequence.
- 10. A transposable element isolatable by the method of Claim 6, 7, 8 or 9.

- -- 36 --
- 11. A method of isolating activation sequences comprising:
- (a) inserting a marker gene within the inverted repeats of a transposable element of Claim 1 to form a modified marker gene having the structure IR-marker-IR;
 - (b) inserting the modified marker gene into a DNA target:
 - (c) selecting for expression of the modified marker; and
 - (d) isolating DNA upstream of said modified marker gene in said DNA target, which DNA upstream of said modified marker gene comprises an activation sequence driving expression of said modified marker gene.
- 12. A method for inactivating a gene in a host cell wherein said gene encodes a gene product, the method comprising:
- (a)transforming a host cell with a genetic element to create a transformed host cell, wherein the genetic element comprises DNA for the gene and a transposable element of Claim 1 inserted within the DNA; and
 - (b)selecting for the transformed host cells which are deficient in the gene product.
- 13. A method for activating a desired gene in a host cell, the method comprising:

 (a)inserting a regulatory gene within the inverted repeats of a transposable element of Claim 1 to form a modified regulatory gene having the structure IR-regulatory gene-IR;
- (b)inserting the modified regulatory gene in DNA comprising the desired gene to form a DNA construct containing the modified regulatory gene upstream of said desired gene;
 - (c)transforming the host cell with the DNA construct; and
 - (d)selecting for transformants having enhanced expression of said desired gene.
- 14. A transposable element isolated from *Aspergillus oryzae* comprising a DNA fragment of at least about 1.2 kb.
- 15. The transposable element of Claim 14 comprising an inverted repeat DNA sequence of SEQ ID NO:17 or a variation thereof.
- 16. A fragment of the transposable element of 14 comprising part of all of the DNA sequence of SEQ ID NO:17 or a variation thereof.

- 17. A method of isolating a transposable element from a filamentous fungus, comprising the steps of:
 - (a) hybridizing fungal DNA under low stringency conditions to a probe, wherein the probe comprises part or all of one of the DNA fragments of Claim 16; and
 - (b) isolating fungal DNA which hybridizes to said probe.
- 18. A method of isolating activation sequences comprising:
- (a) inserting a marker gene within the inverted repeats of a transposable element of Claim 10 to form a modified marker gene having the structure IR-marker-IR;
 - (b) inserting the modified marker gene into a DNA target;
 - (c) selecting for expression of the modified marker; and
 - (d) isolating DNA upstream of said modified marker gene in said DNA target, which DNA upstream of said modified marker gene comprises an activation sequence driving expression of said modified marker gene.
- 19. A method for inactivating a gene in a host cell wherein said gene encodes a gene product, the method comprising:
- (a) transforming a host cell with a genetic element to create a transformed host cell, wherein the genetic element comprises DNA for the gene and a transposable element of Claim 10 inserted within the DNA; and
 - (b) selecting for the transformed host cells which are deficient in the gene product.
- 20. A method for activating a desired gene in a host cell, the method comprising:
- (a) inserting a regulatory gene within the inverted repeats of a transposable element of Claim 10 to form a modified regulatory gene having the structure IR-regulatory gene-IR;
- (b) inserting the modified regulatory gene in DNA comprising the desired gene to form a DNA construct containing the modified regulatory gene upstream of said desired gene;
 - (c) transforming the host cell with the DNA construct; and
- (d)selecting for transformants having enhanced expression of said desired gene.

1/13



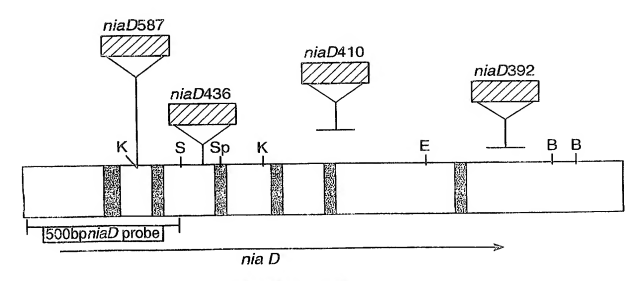
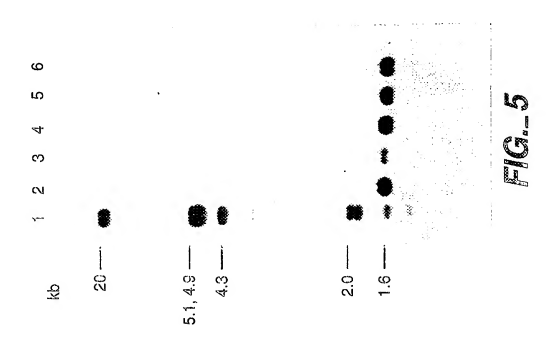
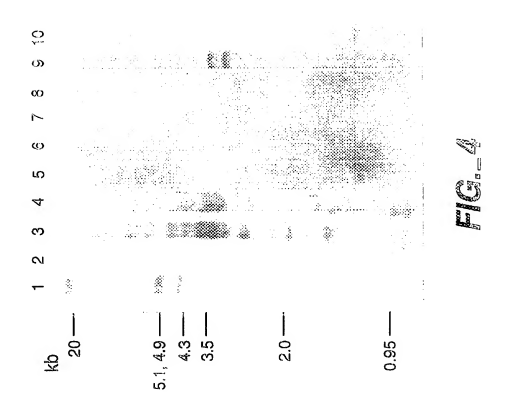
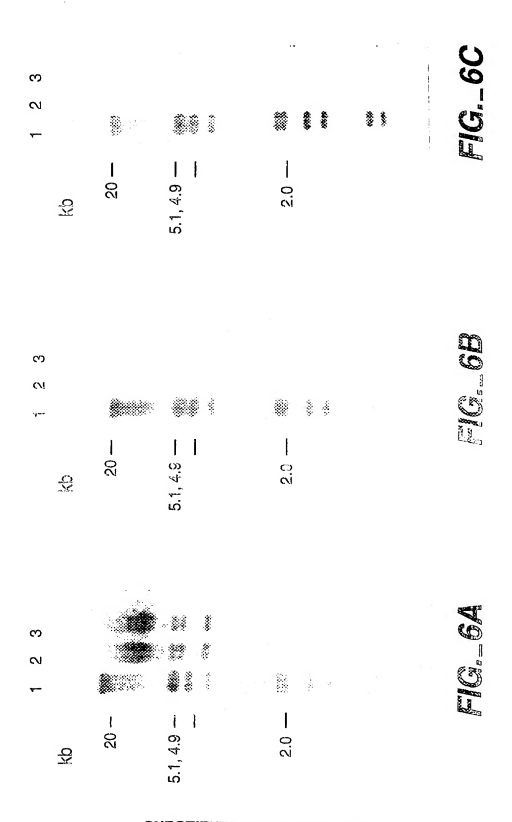


FIG. 2
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tgtcgacggctctctggactggcccaatgatggcagatatcctacggagtgcga
agcctttgaggaaagccaagtaACGTAATCAA CGGTCGAACG GGCCACACGG

TCAGGCGGGC CATCCTGAAA TCCCATATAA AAGATGTCTT GGGGATTCTA
TTATATATCA ACCAGTACTA CTTCTATGAA GCTCTAACTT TGTAGATAGT
TATATATATA AGAATAAGTA TTCCATGAAT TTTTCAGATT TTAGAATTTT
TACTTTGATA ATGAAACCAG ATTCTTATAT AAAACATATA AATACAGATA
TTGTAATATG ATAAGTCCAT AAGTAAAAGT ATATTCATTT TTAGAAGGTA
TATAGATATT ATTTATATTA TTTAAAATCT ATATAGAAGA AATCTAATTC
TTCTAGACCT GGATGGTAGA GATATATTAT GTTTAAAAAG ATATCTTTTG
TATAGTATTA CCAGATGGCC CGCCTGACCG TGTGGCCCGT CCGACCGTTG
ATTACGTtatgccttgatggaggagctgataagctggtaagttaccttatcca
tccatqcatgcatgccctga

FIG._7

099

CTACATATAA T Y K

GGAATGATTG G M I A

ATGAAACAGG E T G

TACAATTTCG Y N F D

AGATGATATA D D I

601

009

GGATCTCATC I S S

CAAGAGTACG (Q E Y G

GGAAGTTATT E V I

GACTGGTTCC AACGCGTACA D W F Q R V Q

AGTTATAAAA V I K

| 60 | 180 | 240 | 300 | 360 | 420 | 480 | 540 |
|---|--|--|---|--|--|--|--|
| AAAACACCAC CATCTATCCC S I P | CTATTCAGAA I Q K | GAACTACTCT T T L | GTCAAAAATT Q K L | AGCGAGGAGC R G A | AACGAGGTTA R G Y | GCCACGAATC H E S | AGGATCCTGA D P E |
| ACGTAATCAA CGGTCGGACG GGCCACACGG TCAGGCGGGC CATCCCTTCG AAAACACCAC CTTGAATCAC CTACCCGAGG CTTTTCAACC ACCACAAATG CCACCAAAAG CATCTATCCC M P P K A S I P | ATCAAAATCG CAGGTGGAGGCAG GATTCTTCTT GCCATTGAAG CTATTCAGAAS ${ m S}$ ${ m K}$ ${ m S}$ ${ m Q}$ ${ m V}$ ${ m E}$ ${ m Q}$ ${ m E}$ ${ m G}$ ${ m E}$ ${ m E}$ ${ m G}$ ${ m E}$ ${ m E$ | AGGCCAAATC ACTAGTATTC GTGAAGCAGC GCGTGTTTAT GACGTCGCTC GAACTACTCT G Q I T S I R E A A R V Y D V A R T T L | CCAGGCTCGA TTATCTGGAC GTGTTTTCGC TAAAATATG ACCAACGCAC GTCAAAAATT Q A R L S G R V F A K N M T N A R Q K L | GTCAAATAAT GAAGAGGAAT CGCTTGTTAA ATGGATCCTA TCTCTAGATA AGCGAGGAGC S N N E E E S L V K W I L S L D K R G A | AAGCCCCCGG CCACTTGATA TCAGAGATAT GGCTAATTTG ATTATCTCTA AACGAGGTTA S P R P L D I R D M A N L I I S K R G Y | TTCAACTGTT GAACAAGTAG GCATCAACTG GGCTTATAGC TTTGTTAAAC GCCACGAATC S T V E Q V G I N W A Y S F V K R H E S | CCTACGAACT CGATTTGCTA GACGACTCAA CTATCCAAGA GCTAAAATGG AGGATCCTGA L R T R F A R L N Y P R A K M E D P E |
| TCAGGCGGGC ACCACAAATG M | GATTCTTCTT I L L | GCGTGTTTAT R V Y | TAAAAATATG K N M | ATGGATCCTA W I L | GGCTAATTTG A N L | GGCTTATAGC A Y S | CTATCCAAGA Y P R |
| GGCCACACGG CTTTTCAACC | AGGAAGGCAG E G R | GTGAAGCAGC E A A | GTGTTTTCGC V F A | CGCTTGTTAA L V K | TCAGAGATAT R D M | GCATCAACTG I N W | GACGACTCAA R L N |
| CGGTCGGACG | CAGGTGGAGC Q V E Q | ACTAGTATTC T S I R | TTATCTGGAC L S G R | GAAGAGGAAT E E E S | CCACTTGATA P L D I | GAACAAGTAG E Q V G | CGATTTGCTA R F A R |
| ACGTAATCAA CTTGAATCAC | ATCAAAATCG S K S | AGGCCAAATC G Q I | CCAGGCTCGA Q A R | GTCAAATAAT S N N | AAGCCCCCGG S P R | TTCAACTGTT S T V | CCTACGAACT L R T |
| 1 | 121 | 181 | 241 | 301 | 361 | 421 | 481 |

FIG._84

| PCT/US97/14978 |
|----------------|
| |

1200

1260

ATCAAAGTCG AATATAATCA GTGGTTTTAG AGCAACAGGT CTTGTTCCTC TAGATCCTGA S K S N I I S G F R A T G L V P L D P D

CAATCATATC GACAAGCTTG ACTTCTTAGA GGTCTATCCT AAAGCTCACC AGTGTGCTTT N H I D K L D F L E V Y P K A H Q C A L

| | | | | 6 | / 13 | | |
|--|---|---|---|---|---|---|---|
| 720 | 780 | 840 | 006 | 096 | 1020 | 1080 | 1140 |
| GGAATCGGGA | CGACCCTGAT | CTCCTACCTG | GATGGCTTCC | TCCTAGTCCT | AAAATGAGGT | ATGTTGGTTG | AATACGGCCG |
| N R E | T L I | P T W | W L P | L V L | N E V | V G C | Y G R |
| GA GGGCAGGTCG GCCGTCCCTA GTTCAACCAG GGAATCGGGA | ATGGGTCACT CCAATTGAGT GTATTCGCTC TAATGGAGAG GTTCTACCTT CGACCCTGAT | CTTTAAAGGC AAAACACATC TAAAGGCATG GTATGAAGGT CAATCTATTC CTCCTACCTG | GAGATTTGAA GTCAGTGATA ATGGTTGGAC TACTGATAAA ATTGGACTTC GATGGCTTCC | AAAACACTTC ATTCCCTTGA TTAGAGGCAA ATCAGTAGGC AAATATAGCC TCCTAGTCCT | CGATGGCCAC GGTAGTCATT TGACACCTGA ATTCGACCAA TCCTGTGCTG AAAATGAGGT | TATACCTATT TGTATGCCAG CTCATTCGTC CCATCTACTT CAGCCTCTTG ATGTTGGTTG | TTTTAGTGTG CTTAAACGCA CGTACGGAGG CATGGTTCCC AAGCAGATGC AATACGGCCG |
| R A G R P S L V Q P G N R E | W V T P I E C I R S N G E V L P S T L I | F K G K T H L K A W Y E G Q S I P P T W | R F E V S D N G W T T D K I G L R W L P | K H F I P L I R G K S V G K Y S L L V L | D G H G S H L T P E F D Q S C A E N E V | I P I C M P A H S S H L L Q P L D V G C | F S V L K R T Y G G M V P K Q M Q Y G R |
| GCCGTCCCTA | TAATGGAGAG | GTATGAAGGT | TACTGATAAA | ATCAGTAGGC | ATTCGACCAA | CCATCTACTT | CATGGTTCCC |
| P S L | N G E | Y E G | T D K | S V G | F D Q | H L L | M V P |
| GGGCAGGTCG | GTATTCGCTC | TAAAGGCATG | ATGGTTGGAC | TTAGAGGCAA | TGACACCTGA | CTCATTCGTC | CGTACGGAGG |
| A G R | I R S | K A W | G W T | R G K | T P E | H S S | Y G G |
| AGTTCCCAGA | CCAATTGAGT | AAAACACATC | GTCAGTGATA | ATTCCCTTGA | GGTAGTCATT | TGTATGCCAG | CTTAAACGCA |
| S S Q R | P I E C | K T H L | V S D N | I P L I | G S H L | C M P A | L K R T |
| AGTAGTAACT AGTTCCCA | ATGGGTCACT | CTTTAAAGGC | GAGATTTGAA | AAAACACTTC | CGATGGCCAC | TATACCTATT | TTTTAGTGTG |
| V V T S S Q | W V T | F K G | R F E | K H F | D G H | I P I | F S V |
| 661 | 721 | 781 | 841 | 901 | 961 | 1021 | 1081 |
| | | | | SUBST | ITUTE S | HEET (R | ULE 26) |

FIG._8B

1201

| 1320 | 1380 | 1440 | 1500 | 1560 | 1620 | 1680 | 1740 | 1800 |
|---|--|--|--|--|--|---|---|--|
| TCAAGTGCTT TCTCGACTCC ATATTCGCTT GAAAACACCA CCAACCCGG ATAGCCAGTC Q V L S R L H I R L K T P P T P D S Q S | AAGTGGCTCA GTGCTTCAAA CACCACATAA TATAAAACAC CTTTTGGAGC ATCCAAAATC S G S V L Q T P H N I K H L L E H P K S | AGTGGAACGC CTACTTCGGA AACGGCAAGC AAGTCCAACT TCACCTACAA ACTCTACACT V E R L L R K R Q A S P T S P T N S T L | ACGTCAGCTT CTCAAAGGGT GTGAACTAGC AATAACAAAC TCAATCATAC TGGCTAAGGA R Q L L K G C E L A I T N S I I L A K E | GAATGCGGAA TTACGTGCTA GCCATGAAAA GCAACTACCA AAGAGGAAGC GTTCAAGGAA N A E L R A S H E K Q L P K R K R S R K | GCAGGTGATC TATACAGAAG GCACTACCGT TGAAGAGGCC CAGAGAGCTA TACAGGAAGT Q V I Y T E G T T V E E A Q R A I Q E V | GGAAGAGGTG CAGAATGATG AAGATATTGA GGTTGAACCC CAATCTCAAT ATACGGAGAC ${ m E}$ E ${ m E}$ V ${ m E}$ P ${ m Q}$ S ${ m Q}$ Y T ${ m E}$ T | CCCCTCGCGC GCGCCTCCAC GCTGCAGTAA TTGCTTCAAT ATAGGCCACC GACGTACACA | GTGTTCTAAA CCACCTACTA ATTAGTTAGA TAGCTGTTTT TACAAGCATT TATGTTGATT C S K P P T N * |
| CCAACCO | CTTTTG | TCACCTY | TCAATCZ | AAGAGGZ | CAGAGA(| CAATCT(| ATAGGC(| TACAAGO |
| P T | L L | S P | S I | K R | Q R | Q S (| I G] | |
| GAAAACACCA | TATAAAACAC | AAGTCCAACT | AATAACAAAC | GCAACTACCA | TGAAGAGGCC | GGTTGAACCC | TTGCTTCAAT | TAGCTGTTTT |
| K T P | I K H | S P T | I T N | Q L P | E E A | V E P | C F N | |
| ATATTCGCTT | CACCACATAA | AACGGCAAGC | GTGAACTAGC | GCCATGAAAA | GCACTACCGT | AAGATATTGA | GCTGCAGTAA | ATTAGTTAGA |
| I R L | P H N | R Q A | E L A | H E K | T T V | D I E | C S N | * |
| TCTCGACTCC | GTGCTTCAAA | CTACTTCGGA | CTCAAAGGGT | TTACGTGCTA | TATACAGAAG | CAGAATGATG | GCGCCTCCAC | CCACCTACTA |
| S R L H | V L Q T | L L R K | L K G C | L R A S | Y T E G | Q N D E | A P P R | P P T N |
| | AAGTGGCTCA | AGTGGAACGC | ACGTCAGCTT | GAATGCGGAA | GCAGGTGATC | GGAAGAGGTG | CCCCTCGCGC | GTGTTCTAAA |
| | S G S | V E R | R Q L | N A E | Q V I | E E V | P S R | C S K |
| 1261 | 1321 | 1381 | 1441 | 1501 | 1561 | 1621 | 1681 | 1741 |
| | | | | | | | | |

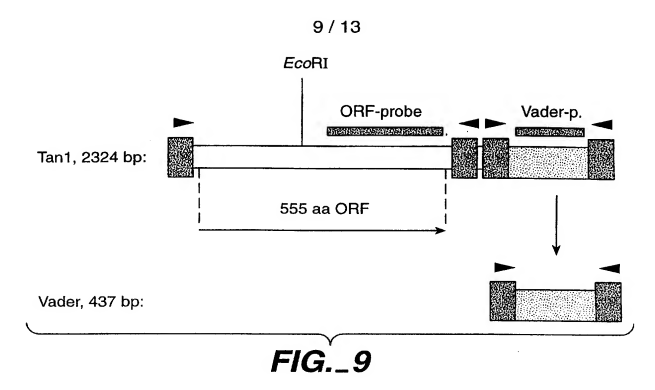
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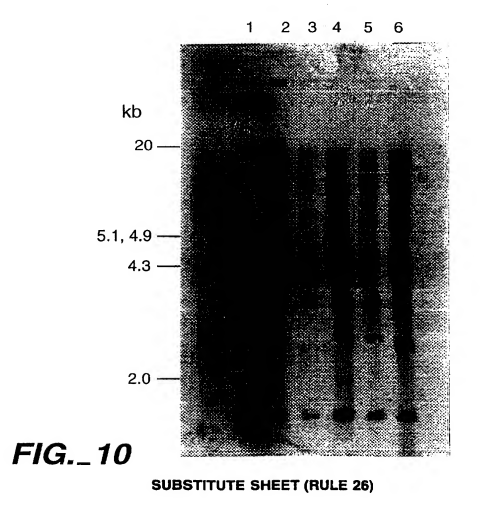
| 1860 1915 | 1975 | 2035 | 2095 | 2155 | 2215 | 2275 | 2320 |
|---|----------------------------------|----------------------------------|-----------------------|----------------------------------|---|--|---|
| CCTGACCGTG | ATATATCTCT | TATAAATAAT | ATTACAATAT | ATTCTAAAAT | TTAGAGCTTC | TATGGGATTT | |
| TCCTACCGGG AGATGGCCCG CCTGACCGTG ACGTAATCAA CGGTCGGACG GGCCCCCGG | AAGATATCTT TTTAAACATA ATATATCTCT | TCTATATAGA TTTTAAATAA TATAAATAAT | GACTTATCAT ATTACAATAT | GTTTCATTAT CAAAGTAAAA ATTCTAAAAT | TTATTCTTAT ATATAACT ATCTACAAAG TTAGAGCTTC | TATATATAG AATCCCCAAG ACATCTTTTA TATGGGATTT | TACGT |
| TCCTACCGGG ACGTAATCAA | AAGATATCTT | TCTATATAGA | TTTACTTATG | GTTTCATTAT | ATATATAACT | AATCCCCAAG | GACCGTTGAT |
| TAGAGGCCTC ATTTGGATCA TATCGGGTAA TCCTACCGGG AGATGGCCCG CCTGACCGTG TGGCCCGCCC GACCGTTGAT TACGTNNNNN ACGTAATCAA CGGTCGGACG GGCCCCCGG | TACTATACCA | | TGAATATACT | TAAGAATCTG | TTATTCTTAT | | CAGGATGGCC GCCGACCGTG TGGCCCGTCC GACCGTTGAT TACGT |
| TAGAGGCCTC ATTTGGATCA TATCGGGTAA TGGCCCGCCC GACCGTTGAT TACGTNNNNN | TCCGGCGGGC CATCTGGTAA TACTATACCA | TCTAGGAGAA | CTTCTAAAAA | ATGTATTATA | CTGAAAATT CATGGAATAC | TACTGGTTGA | GCCGACCGTG |
| TAGAGGCCTC | TCCGGCGGGC | ACCATCCAGG | ATCTATATAC | CTGTATTTAT | CTGAAAAATT | ATAGAAGTAG | CAGGATGGCC |
| 1801 1861 | 1916 | 1976 | 2036 | 2096 | 2156 | 2216 | 2276 |

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F/G._8L

WO 98/08960 PCT/US97/14978





10 / 13

| 60 | 180 | 240 | 300 | 360 | 420 | 480 | 540 | 009 | 099 |
|---|--|---|--|---|---|---|---|---|---|
| AAAACACCAC CATCTATCCC S I P | CTATTGAGAA I R K | GAACTACTCT T T L | GTCAAAAATT Q K L | AGCGAGGAGC R G A | AACGAGGTTA R G Y | GCCACGAATC H E S | AGGATCCTGA D P E | GGATCTCATC I S S | CTACATATAA T Y K |
| ACGTAATCAA CGGTCGGGCG GGCCACGG TCAGGCGGGC CACCCCTTCG AAAACACCAC | ATCAAAATCG CAGGTGGAAGGCAG GATTCTTCTT GCCATTGAAG CTATTGAGAA | AGGCCAAATC ACTAGTATTC GTGAAGCAGC GCGTGTTTAT GACGTCGCTC GAACTACTCT | CCAGGCTCGA TTATCTGGAC GTGTTTTCGC TAAAATATG ACCAACGCAC GTCAAAAATT | GTCAAATAAT GAAGAAAT CGCTTGTTAA ATGGATCCTA TCTCTAGATA AGCGAGGAGC | AAGCCCCCGG CCACTTGATA TCAGAGATAT GGCTAATTTG ATTATCTCTA AACGAGGTTA | TTCAACTGTT GAACAAGTAG GCATCAACTG GGCTTATAGC TTTGTTAAAC GCCACGAATC | CCTACGAACT CGATTTGCTA GACGACTCAA CTATCAAAGA GCTAAAATGG AGGATCCTGA | AGTTATAAAA GACTGGTTCA AACGCGTACA GGAAGTTATT CAAGAGTACG GGATCTCATC | AGATGATATA TACAATTTCG ATGAAACAGG GTTTGCTATG GGAATGATTG CTACATATAA |
| CTTGAATCAC CTACCCGAGG CTTTTCAACC ACCACAAATG CCACCAAAAG CATCTATCCC | S K S Q V E R E G R I L L A I E A I R K | G Q I T S I R E A A R V Y D V A R T T L | Q A R L S G R V F A K N M T N A R Q K L | S N N E E E S L V K W I L S L D K R G A | S P R P L D I R D M A N L I I S K R G Y | S T V E Q V G I N W A Y S F V K R H E S | L R T R F A R R L N Y Q R A K M E D P E | V I K D W F K R V Q E V I Q E Y G I S S | D D I Y N F D E T G F A M G M I A T Y K |
| TCAGGCGGGC ACCACAATG M | GATTCTTCTT I L L | GCGTGTTTAT R V Y | TAAAAATATG K N M | ATGGATCCTA W I L | GGCTAATTTG A N L | GGCTTATAGC A Y S | CTATCAAAGA Y Q R | GGAAGTTATT E V I | GTTTGCTATG F A M |
| GGCCACACGG | GGGAAGGCAG | GTGAAGCAGC | GTGTTTTCGC | CGCTTGTTAA | TCAGAGATAT | GCATCAACTG | GACGACTCAA | AACGCGTACA | ATGAAACAGG |
| CTTTTCAACC | E G R | E A A | V F A | L V K | R D M | I N W | R L N | R V Q | E T G |
| CGGTCGGGCG | CAGGTGGAGC | ACTAGTATTC | TTATCTGGAC | GAAGAGGAAT | CCACTTGATA | GAACAAGTAG | CGATTTGCTA | GACTGGTTCA | TACAATTTCG |
| CTACCCGAGG | Q V E R | T S I R | L S G R | E E E S | P L D I | E Q V G | R F A R | D W F K | Y N F D |
| ACGTAATCAA | ATCAAAATCG | AGGCCAAATC | CCAGGCTCGA | GTCAAATAAT | AAGCCCCCGG | TTCAACTGTT | CCTACGAACT | AGTTATAAAA | AGATGATATA |
| CTTGAATCAC | S K S | G Q I | Q A R | S N N | S P R | S T V | L R T | V I K | D D I |
| 611 | 121 | 181 | 241 | 301 | 361 | 421 | 481 | 541 | 601 |

FIG._11B

| | | | | 11 | / 13 | | | | |
|---|---|---|---|---|---|---|---|---|---|
| 720 | 780 | 840 | 006 | 096 | 1020 | 1080 | 1140 | 1200 | 1260 |
| GGAATCGGGA | CGACCCTGAT | CTCCTACCTG | GATGGCTTCA | TCCTAGTCCT | AAAATGAGGT | ATGTTGGTTG | AATACGGCCG | AGTGTGCTTT | TAGATCCTGA |
| N R E | T L I | P T W | W L Q | L V L | N E V | V G C | Y G R | C A L | D P D |
| AGTAGTAACT AGTTCCCAGA GGGCAGGTCG GCCGTCCCTA GTTCAACCAG GGAATCGGGA | ATGGGTCACT GCAATTGAGT GTATTCGCTC TAATGGAGAG GTTCTACCTT CGACCCTGAT | CTTTAAAGGC AAAACACATC TAAAGGCATG GTATGAAGGT CAATCTATTC CTCCTACCTG | GAGATTTGAA GTCAGTGATA ATGGTTGGAC TACTGATAAA ATTGGACTTC GATGGCTTCA | AAAACACTTC ATTCCCTTGA TTAGAGGCAA ATCAGTAGGC AAATATAGCC TCCTAGTCCT | CGATGGCCAC GGTAGTCATT TGACACCTGA ATTCGACCAA TCCTGTGCTG AAAATGAGGT | TATACCTATT TGTATGCCTG CTCATTCGTC CCATCTACTT CAGCCTCTTG ATGTTGGTTG | TTTTAGTGTG CTTAAACGCA CGTACGGAGG CATGGTTCAA AAGCAGATGC AATACGGCCG | CAATCATATC GACAAGCTTG ACTTCTTAGA GGTCTATCCT AAAGCTCACC AGTGTGCTTT | ATCAAAGTCG AATATAATCA GTGGTTTTAG AGCAACAGGT CTTGTTCCTC TAGATCCTGA |
| V V T S S Q R A G R P S L V Q P G N R E | W V T A I E C I R S N G E V L P S T L I | F K G K T H L K A W Y E G Q S I P P T W | R F E V S D N G W T T D K I G L R W L Q | K H F I P L I R G K S V G K Y S L L V L | D G H G S H L T P E F D Q S C A E N E V | I P I C M P A H S S H L L Q P L D V G C | F S V L K R T Y G G M V Q K Q M Q Y G R | N H I D K L D F L E V Y P K A H Q C A L | S K S N I I S G F R A T G L V P L D P D |
| GCCGTCCCTA | TAATGGAGAG | GTATGAAGGT | TACTGATAAA | ATCAGTAGGC | ATTCGACCAA | CCATCTACTT | CATGGTTCAA | GGTCTATCCT | AGCAACAGGT |
| P S L | N G E | Y E G | T D K | S V G | F D Q | H L L | M V Q | V Y P | A T G |
| GGGCAGGTCG | GTATTCGCTC | TAAAGGCATG | ATGGTTGGAC | TTAGAGGCAA | TGACACCTGA | CTCATTCGTC | CGTACGGAGG | ACTTCTTAGA | GTGGTTTTAG |
| A G R | I R S | K A W | G W T | R G K | T P E | H S S | Y G G | F L E | G F R |
| AGTTCCCAGA | GCAATTGAGT | AAAACACATC | GTCAGTGATA | ATTCCCTTGA | GGTAGTCATT | TGTATGCCTG | CTTAAACGCA | GACAAGCTTG | AATATAATCA |
| S S Q R | A I E C | K T H L | V S D N | I P L I | G S H L | C M P A | L K R T | D K L D | N I I S |
| AGTAGTAACT | ATGGGTCACT | CTTTAAAGGC | GAGATTTGAA | AAAACACTTC | CGATGGCCAC | TATACCTATT | TTTTAGTGTG | CAATCATATC | ATCAAAGTCG |
| V V T | W V T | F K G | R F E | K H F | D G H | I P I | F S V | N H I | S K S |
| 661 | 721 | 781 | 841 | 901 | 961 | 1021 | 1081 | 1141 | 1201 |

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GTGTTCTAAA CCACCTACTA ATTAGTTAGA TAGCTGTTTT TACAAGCATT TATGTTGATT C S K P P T N *

| 1261 | TCAAGTGCTT Q V L | TCTCGACTCC S R L H | ATATTCGCTT I R L | GAAAACACCA K T P TATAAAACAC | TCAAGTGCTT TCTCGACTCC ATATTCGCTT GAAAACACCA CCAACCCCGG ATAGCCAGTC Q V L S R L H I R L K T P P T P D S Q S AACACCTCA GTGCTTCAAA CACACATAA TATAAAACAC CTTTTGAAGC ATCCAAAATC | 1320 |
|------|---------------------|----------------------------------|---------------------|-----------------------------------|--|------|
| 1381 | AGTGGAACGC | V L Q T CTACTTCGGA L L R K | AACGGCAAGC R Q A | I K H AAGTCCAACT S P T | AGTGGAACGC CTACTTCGGA AACGGCAAGC AAGTCCAACT TCACCTACAA ACTCTACACT V S P T S P T N S T L | 1440 |
| 1441 | ACGTCAGCTT R Q L | CTCAAAGGGT L K G C | GTGAACTAGC E L A | AATAACAAAC I T N | ACGTCAGCTT CTCAAAGGGT GTGAACTAGC AATAACAAAC TCAATCATAC TGGCTAAGGA R Q L L K G C E L A I T N S I I L A K E | 1500 |
| 1501 | GAATGCGGAA N A E | TTACGTGCTA L R A S | GCCATGAAAA H E K | GCAACTACCA Q L P | GAATGCGGAA TTACGTGCTA GCAACTACCA AAGAGGAAGC GTTCAAGGAA N A E L R A S H E K Q L P K R K R S R K | 1560 |
| 1561 | GCAGGTGATC Q V I | TATACAGAAG Y T E G | GCACTACCGT T T V | TGAAGAGGCC E E A | GCAGGTGATC TATACAGAAG GCACTACCGT TGAAGAGGCC CAGAGAGCTA TACAGGAAGT Q V I Y T E G T T V E E A Q R A I Q E V | 1620 |
| 1621 | GGAAGAGGTG E E V | CAGAATGATG Q N D E | AAGATATTGA D I E | GGTTGAACCC V E P | GGAAGAGGTG CAGAATGATGATTTGA GGTTGAACCC CAATCTCAAT ATACGGAGACEE ${ m E}$ ${$ | 1680 |
| 1681 | CCCCTCGCGC P S R | GCGCCTCCAC A P P R | GCTGCAGTAA CSN | TTGCTTCAAT C F N | CCCCTCGCGC GCGCCTCCAC GCTGCAGTAA TTGCTTCAAT ATAGGCCACC GACGTACACA | 1740 |

FIG._ 11(

| 1860 | 1915 | 1975 | 2035 | 2095 | 2155 | 2215 | 2275 | 2324 |
|----------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|-----------------------|-----------------------|-----------------------|--|
| CCTGACCGTG | 992222229 | ATATATCTCT | TATAAATAAT | ATTACAATAT | ATTCTAAAAA | AGTTAGAGCT | TATATGGGAT | |
| TCCTACCGAG AGATGGCCCG CCTGACCGTG | ACGTAATCAA CGGTCGGACG GGCCCCCCGG | AAGATATCTT TTTAAACATA ATATATCTCT | TCTATATAGA TTTTAAATAA TATAAATAAT | TTTACTTATG GACTTATCAT ATTACAATAT | GTTTCATTAT CAAAGTAAAA | | | TGATTACGT |
| TCCTACCGAG | ACGTAATCAA | AAGATATCTT | TCTATATAGA | TTTACTTATG | GTTTCATTAT | TATATATAAA CTATCTACAA | AGAATCAAAA AGACATCTTT | CCCGCCTGAC CGTGTGGCCC GTTCGACCGT TGATTACGT |
| NTCA TATCGGGTAA | | TACTATACAA | TTAGATTTCT | TGAATATACT | TAAGAATCTG | CTTATTCTTA | GATATATAAT | CGTGTGGCCC |
| ATTTTG | GACCGTTGAT TACGT????? | TCAGGCGGC CATCTGGTAA TACTATACAA | TCTAGGAGAA TTAGATTTCT | CTTCTAAAAA | ATGTATTATA | TCATGGAATA | AGTACTGGTT | CCCGCCTGAC |
| TAGAGGCCTC | TGGCCCGCCC | TCAGGCGGGC | ACCATCCAGG | ATCTATATAC | CTGTATTTAT | TCTGAAAAAT | TCATAGAAGT | TTCAGGATGG |
| 1801 | 1861 | 1916 | 1976 | 2036 | 2096 | 2156 | 2216 | 2276 |

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Int ional Application No PCT/US 97/14978

| A. CLASS IPC 6 | iFICATION OF SUBJECT MATTER C12N15/81 C12N9/22 C12Q1 | /68 C12N15/11 | , |
|---------------------------------------|--|--|--|
| According t | o International Patent Classification(IPC) or to both national clas | ssification and IPC | |
| B. FIELDS | SEARCHED | | |
| Minimum de IPC 6 | ocumentation searched (classification system followed by classi CO7K C12N C12Q | fication symbols) | |
| Documenta | tion searched other than minimum documentation to the extent t | hat such documents are included in the fields se | arched |
| Electronic o | lata base consulted during the international search (name of da | ta base and, where practical, search terms used |) |
| C. DOCUM | ENTS CONSIDERED TO BE RELEVANT | | |
| Category ° | Citation of document, with indication, where appropriate, of the | e relevant passages | Relevant to claim No. |
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| ° Special cat | egories of cited documents : | "T" later document published after the inter or priority date and not in conflict with | national filing date |
| conside "E" earlier d filing da | | cited to understand the principle or the invention "X" document of particular relevance; the c cannot be considered novel or cannot | eory underlying the |
| which i citation | nt which may throw doubts on priority claim(s) or s cited to establish the publicationdate of another or other special reason (as specified) nt referring to an oral disclosure, use, exhibition or neans | involve an inventive step when the doc "Y" document of particular relevance; the cl cannot be considered to involve an inv document is combined with one or mo ments, such combination being obvious | laimed invention ventive step when the re other such docu- |
| later th | nt published prior to the international filing date but an the priority date claimed | in the art. "&" document member of the same patent f | amily |
| | ctual completion of theinternational search December 1997 | Date of mailing of the international sear $30/12/1997$ | rch report |
| | ailing address of the ISA | Authorized officer | |
| | European Patent Office, P.B. 5818 Patentlaan 2 NL – 2280 HV Rijswijk Tel. (+31–70) 340–2040, Tx. 31 651 epo nl, Fax: (+31–70) 340–3016 | Montero Lopez, B | |

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